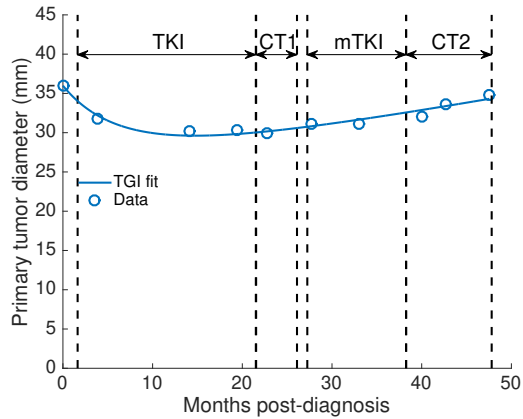
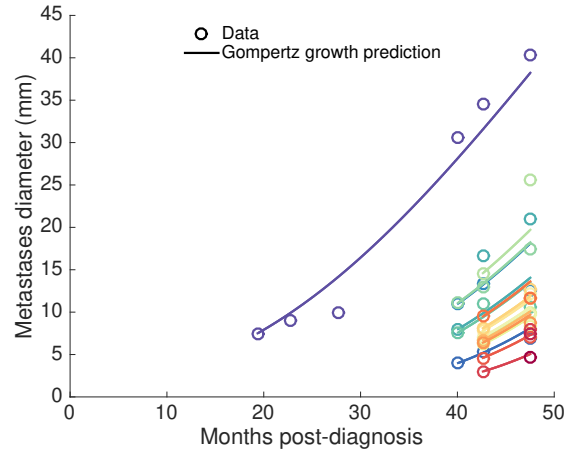


Data of a NSCLC patient with brain mets

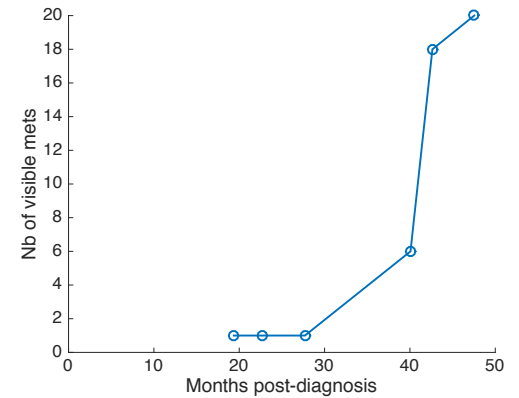
Primary tumor size



Metastases size



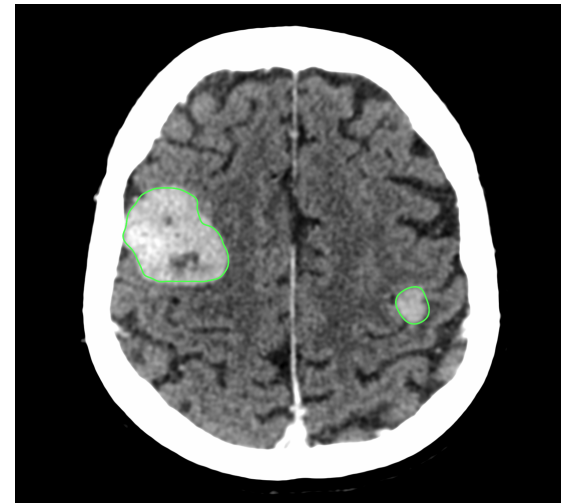
Number of visible mets



Lung CT



Brain CT scan



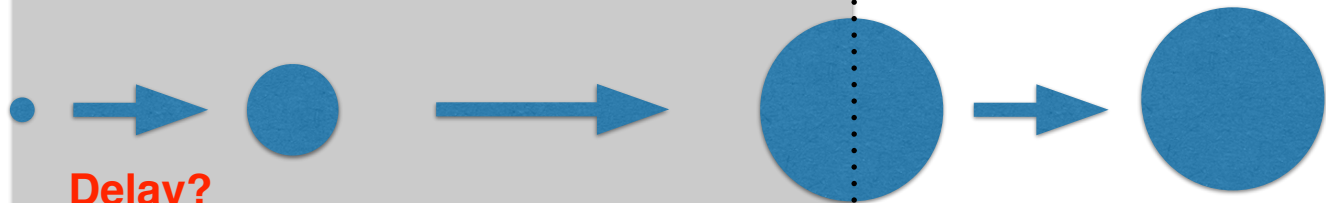
First cancer cell

Diagnosis

treatment

$$\text{Growth law: } g_p(V_p) = V_p(\alpha_p - \beta_p \ln(V_p))$$

Primary Tumor



Delay?

$$\text{Dissemination law: } d(V_p) = \mu(V_p)^\gamma$$

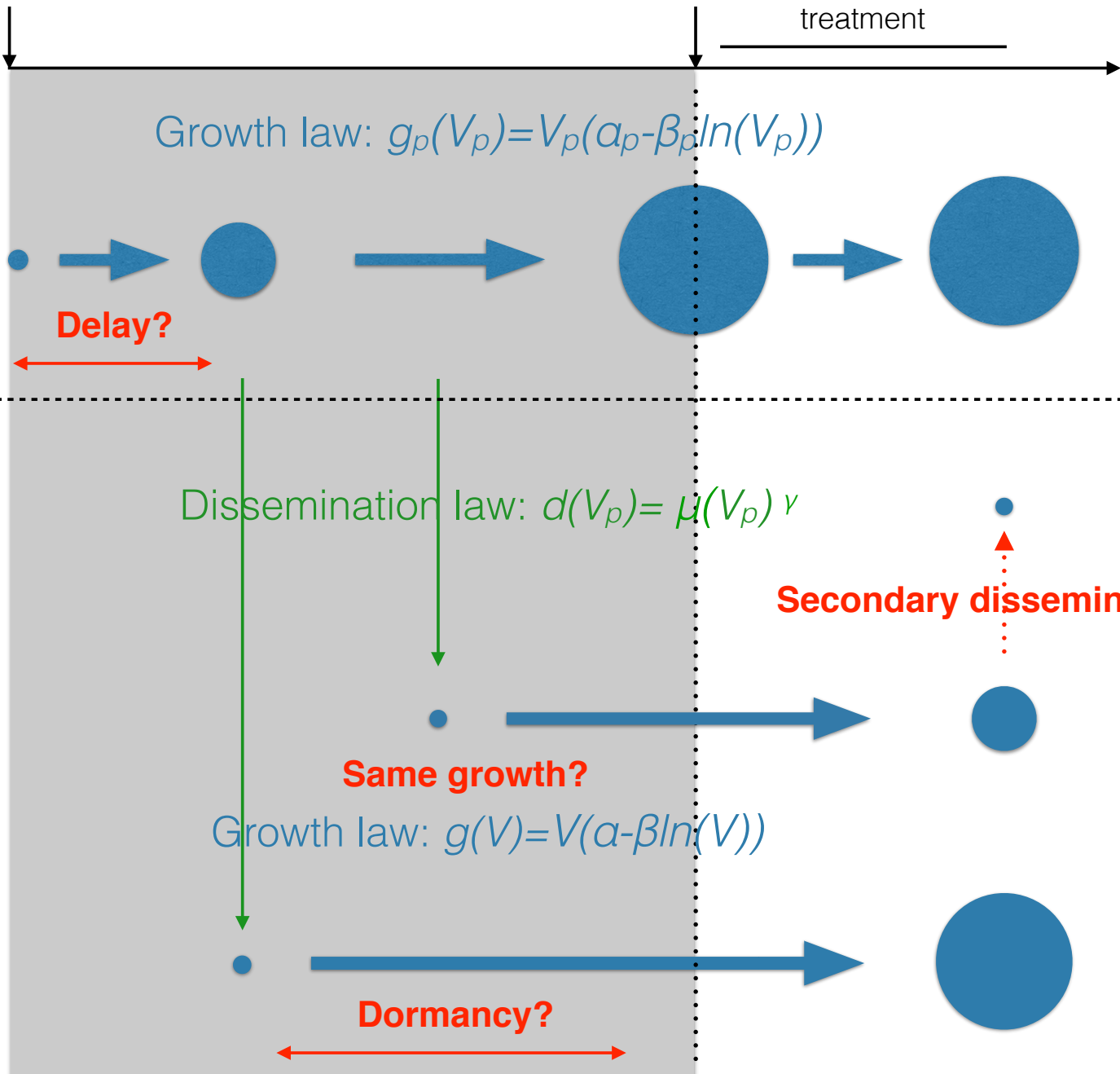
Brain Metastases

Secondary dissemination?

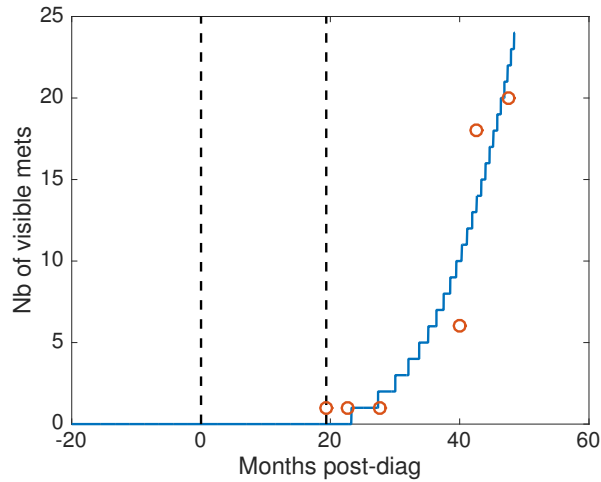
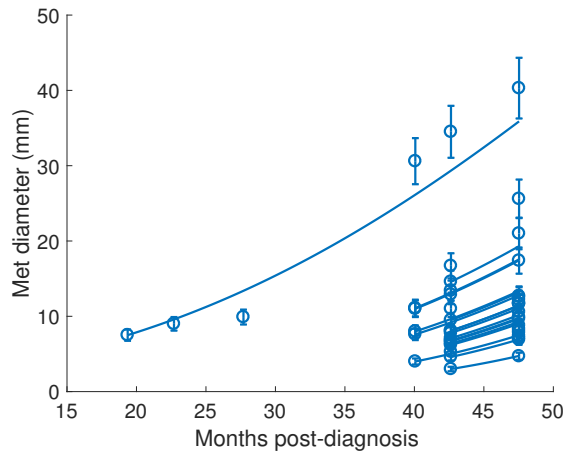
$$\text{Growth law: } g(V) = V(\alpha - \beta \ln(V))$$

Same growth?

Dormancy?



The model with dormancy could describe best the data

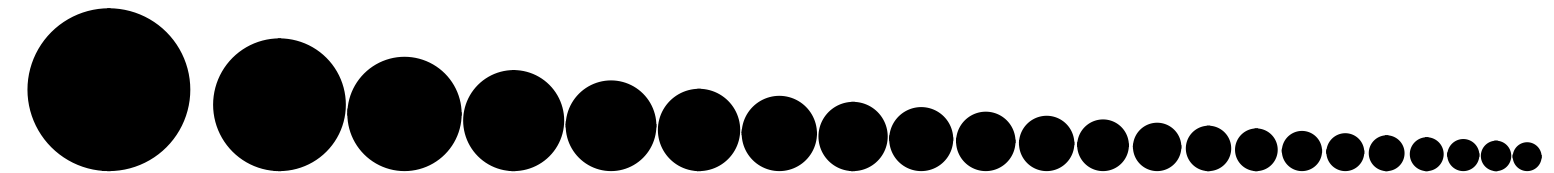


Objective function

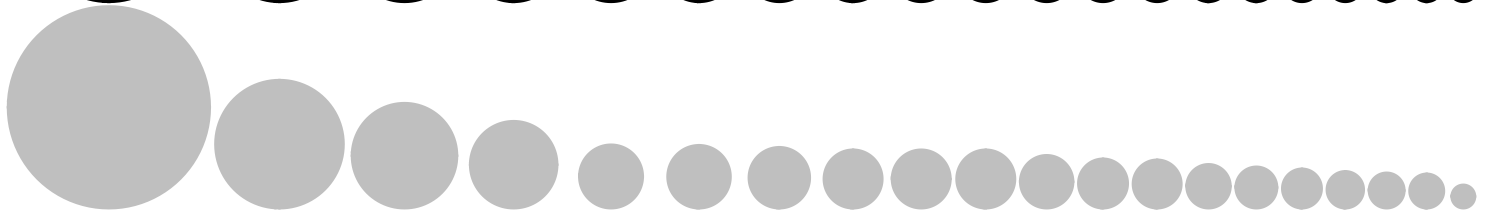
Model	Patient 1	Patient 2
Base	5.51	2.53
Secondary	5.43	2.3
Delay	5.23	1.53
Dormancy	4.93	1.71
Diff. growth	4.95	1.79

Dormancy estimated to 133 days \pm 4.2%

Model



Data



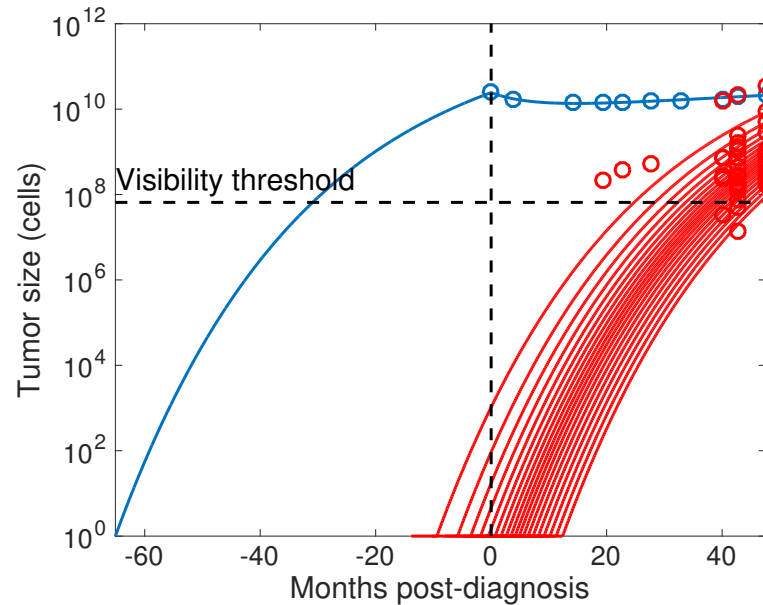
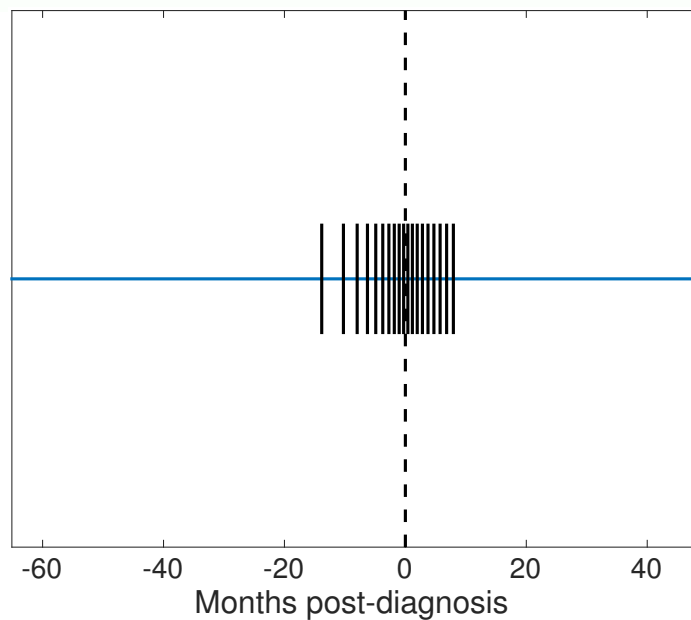
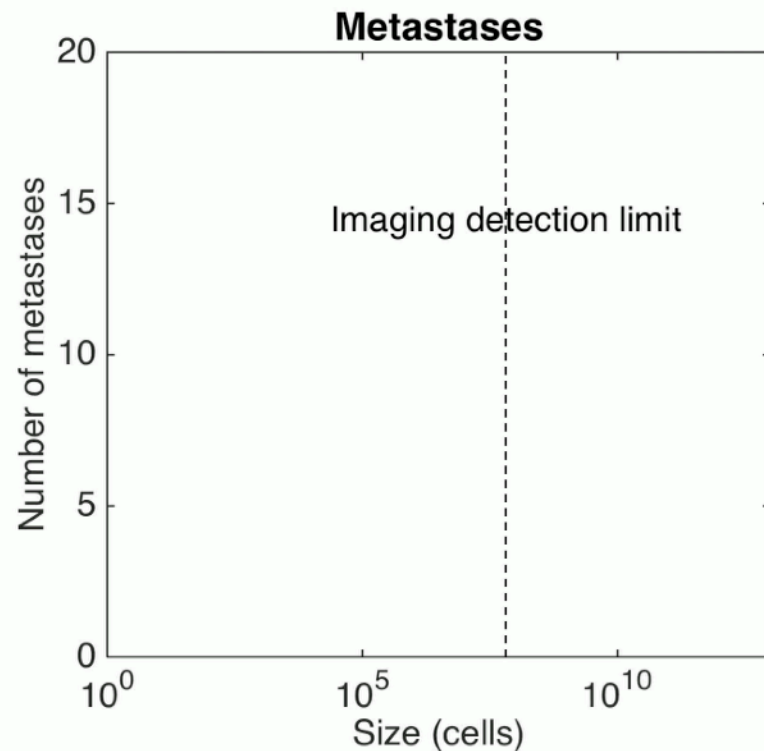
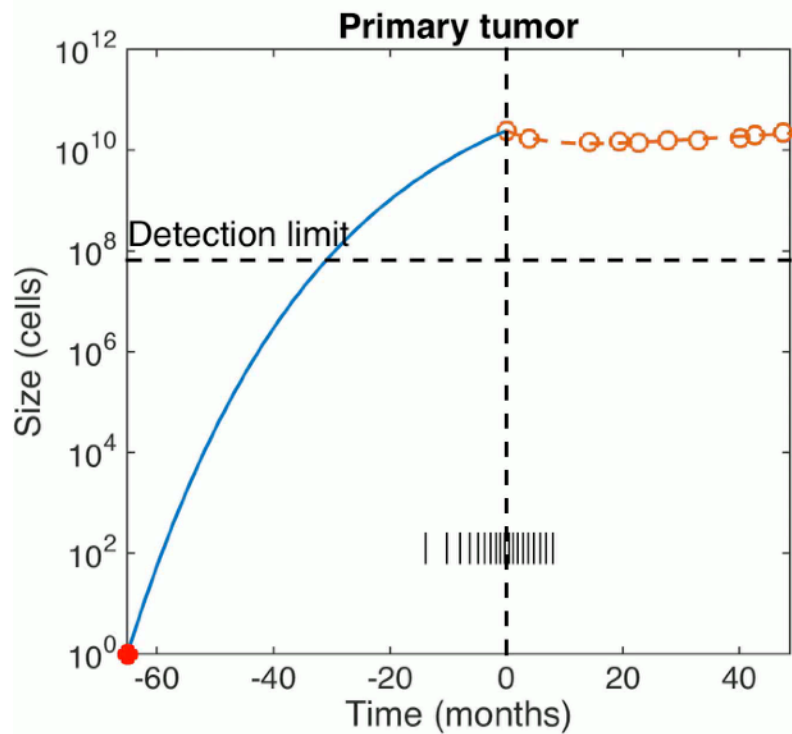
t = -55 months
— 10 mm

*

Primary
tumor
(lung)

Metastases (brain)

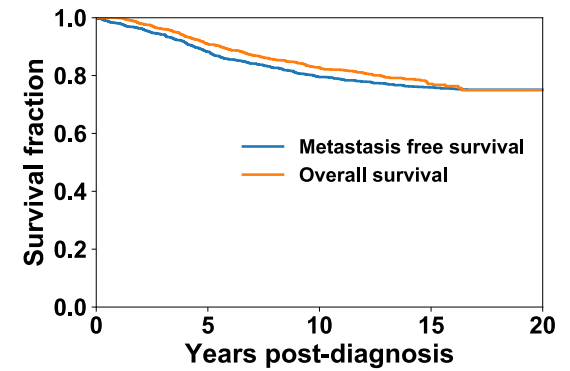
t = -65.1 months



Clinical application - Metastatic relapse in breast cancer

Clinical data of individual breast metastatic relapse

K = 25 features



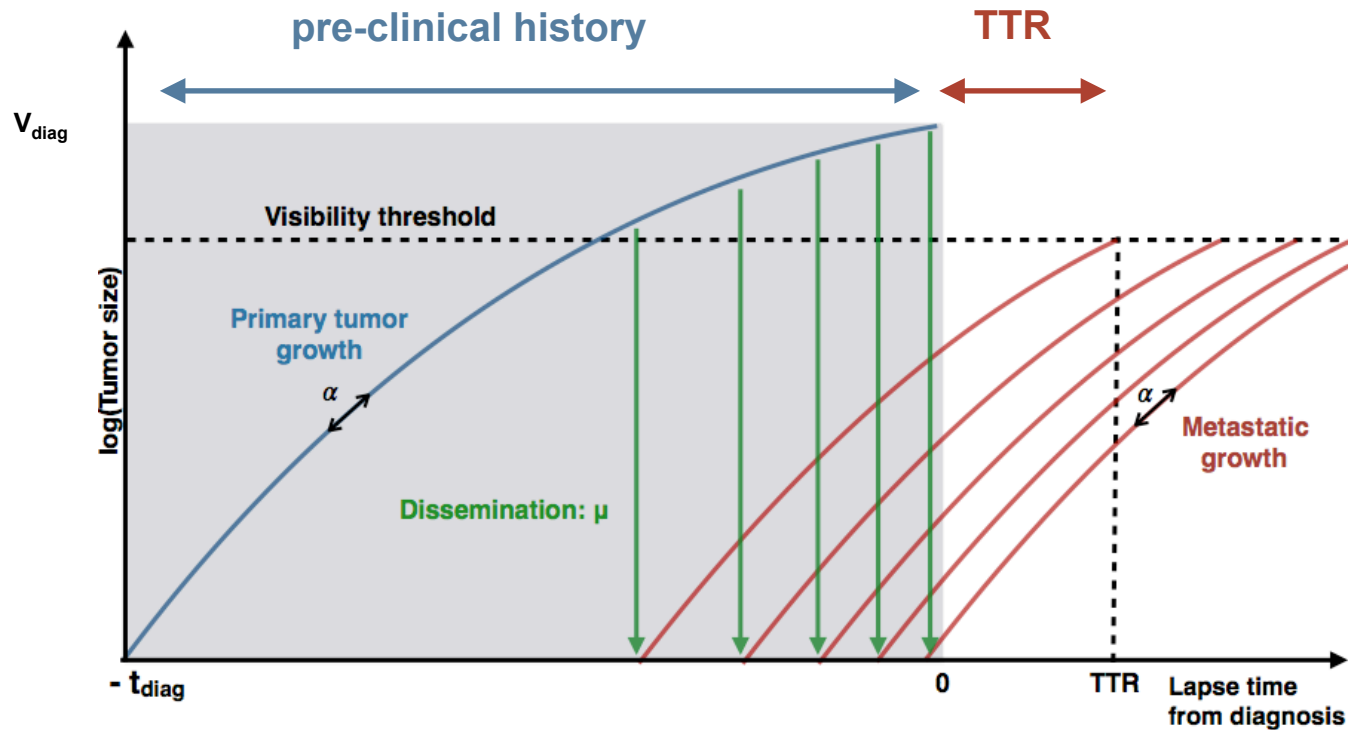
outcome

n = 1057 patients (642 w/o adj)

menopausal_status	ER	PR	Ki67	HER2	HER2_intensity	CK56	EGFR	VIM	ALDH1
Post-ménopause	20	0	0	0	0	0	0	0	0
Ménopause	40	95	8	0	0	0	0	0	0
Activité génitale	87	10	26	0	0	0	0	80	0
Post-ménopause	100	100	8	0	0	0	0	0	0
Post-ménopause	0	0	16	82	+++	0	0	0	0
Activité génitale	100	95	12	0	0	0	0	0	1
Activité génitale	56	100	17	0	0	0	0	0	0
Activité génitale	57	85	23	100	+++	0	0	0	0
Post-ménopause	80	5	20	0	0	0	0	0	0
Post-ménopause	0	0	15	100	+++	0	5	0	0
Post-ménopause	100	80	10	0	0	0	0	0	0
Post-ménopause	30	0	5	0	0	0	0	0	0
Post-ménopause	0	0	15	40	+++	0	0	0	0
Ménopause	0	80	8	0	0	0	0	0	0
Post-ménopause	0	0	27	0	0	0	30	0	1
Post-ménopause	0	0	56	0	0	80	60	100	0
Activité génitale	50	92	2	1	+	0	0	0	0
Post-ménopause	0	47	5	0	0	0	0	80	0
Post-ménopause	65	0	10	0	0	0	0	60	0
Post-ménopause	100	50	11	0	0	0	0	0	0
Ménopause	20	100	0	0	0	0	0	0	0
Activité génitale	90	6	5	0	0	0	0	0	0
Post-ménopause	100	3	5	0	0	0	0	0	0
Activité génitale	0	0	6	0	0	0	0	0	0
Ménopause	80	100	5	0	0	0	0	0	0
Post-ménopause	100	85	25	0	0	0	0	0	0
Post-ménopause	10	45	11	13	+++	0	0	0	0
Post-ménopause	66	1	2	40	++	0	0	0	0

date_metastatic_relapse	date_death_or_loss
854	0
censored	
1999-02-04 00:00:00	1998-04-26 00:00:00
	1999-01-06 00:00:00
	1993-10-21 00:00:00
	2004-06-15 00:00:00
1990-09-04 00:00:00	2006-03-21 00:00:00
1993-02-08 00:00:00	2002-04-05 00:00:00
1999-12-15 00:00:00	2006-11-23 00:00:00
	1997-11-02 00:00:00
	2006-09-15 00:00:00
1995-03-08 00:00:00	2003-03-29 00:00:00
	2003-12-02 00:00:00
1990-04-06 00:00:00	1990-10-20 00:00:00
1994-11-02 00:00:00	2003-10-14 00:00:00
	2004-11-19 00:00:00
	2006-09-30 00:00:00
	1991-07-31 00:00:00
	1995-07-05 00:00:00
	2005-12-08 00:00:00
	2005-05-23 00:00:00
	2007-09-06 00:00:00
	2006-09-06 00:00:00
	2001-02-09 00:00:00
	2005-07-23 00:00:00
	1993-08-12 00:00:00
	1995-01-01 00:00:00
	1993-02-08 00:00:00

Mechanistic modeling of time to relapse



- Number of metastases with size larger than the visible size V_{vis} (= 0.5 cm)

$$N_{vis}(t) = \int_{V_{vis}}^{+\infty} \rho(t, v) dv = \int_0^{t-\tau_{vis}} d(V_p(t)) dt$$

τ_{vis} = time to reach V_{vis}

- Time to relapse (TTR) defined as the time elapsed from diagnosis to the appearance of a first visible metastasis

$$TTR = \inf \{t > 0 : N_{vis}(t_{diag} + t) \geq 1\}$$

- Parameter β fixed such that carrying capacity = 10^{12} cells

Mixed-effects statistical model

$$\ln(T^i) = \ln(TTR(V_{diag}^i; \alpha^i, \mu^i)) + \varepsilon^i, \quad \varepsilon^i \sim \mathcal{N}(0, \sigma^2) \quad (\text{Observation model})$$

$$S(t|\alpha^i, \mu^i) = \mathbb{P}(T^i > t|\alpha^i, \mu^i)$$

Survival function to account for **censoring** in the likelihood

$$\ln(\alpha^i) = \ln(\alpha_{pop}) + \eta_{\alpha}^i, \quad \eta_{\alpha}^i \sim \mathcal{N}(0, \omega_{\alpha}^2)$$

$$\ln(\mu^i) = \ln(\mu_{pop}) + \eta_{\mu}^i, \quad \eta_{\mu}^i \sim \mathcal{N}(0, \omega_{\mu}^2)$$

fixed effects

random effects

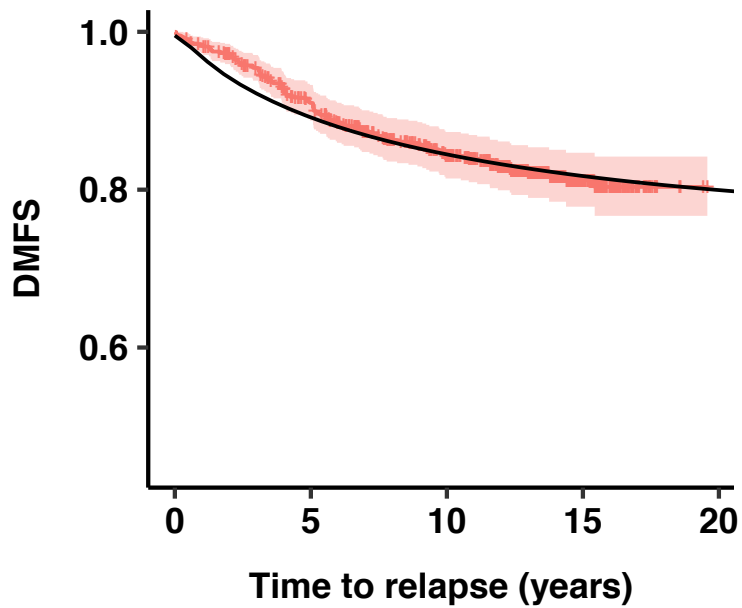


Lavielle, CRC press, 2014

Likelihood maximization performed using the *saemix* R package (SAEM algorithm)

Comets, Lavenu, Lavielle, *J Stat Softw*, 2017

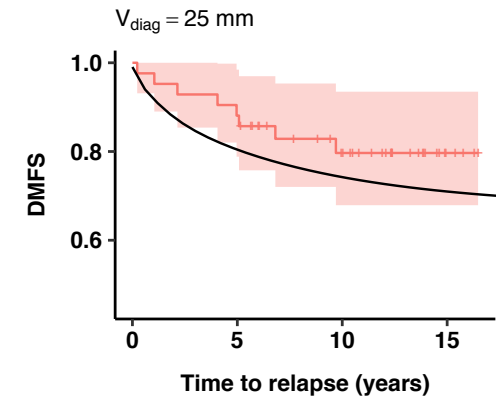
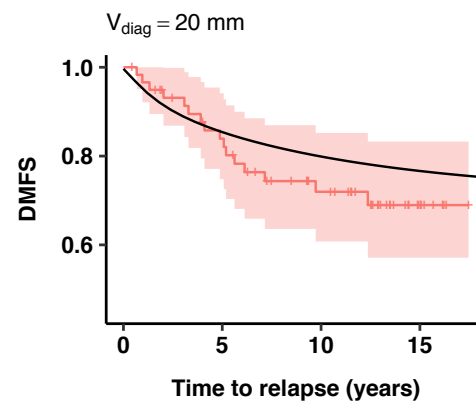
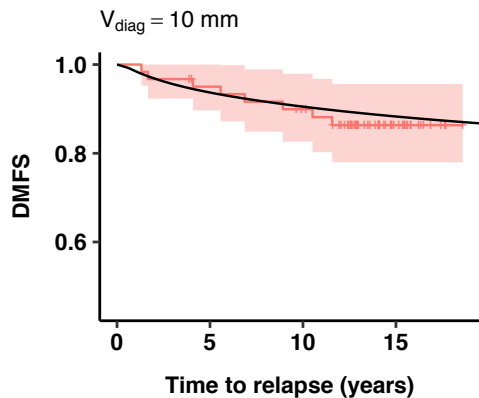
Descriptive power: fit to the data



—+— Kaplan–Meier estimate

— Model fit

Parameter	Estimate	r.s.e. (%)
$\log \alpha_{pop}$	-6.337	12.635
$\log \mu_{pop}$	-26.814	3.683
σ	0.542	28.409
ω_{α}	3.373	36.435
ω_{μ}	3.780	15.876



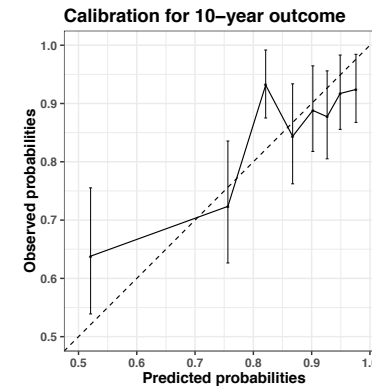
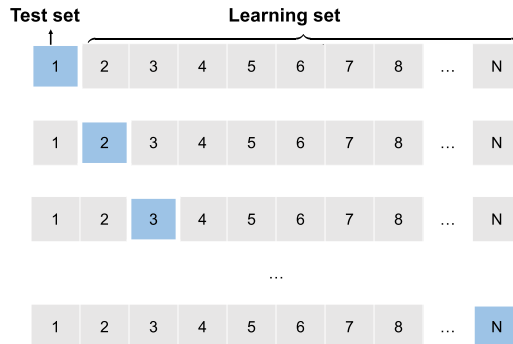
Predictive power: covariates

$$\ln(\mu^i) = \ln(\mu_{pop}) + \beta_{\mu}^T \mathbf{x}_{\mu}^i + \eta_{\mu}^i, \quad \eta_{\mu}^i \sim \mathcal{N}(0, \omega_{\mu}^2)$$

$$\ln(\alpha^i) = \ln(\alpha_{pop}) + \beta_{\alpha}^T \mathbf{x}_{\alpha}^i + \eta_{\alpha}^i, \quad \eta_{\alpha}^i \sim \mathcal{N}(0, \omega_{\alpha}^2)$$

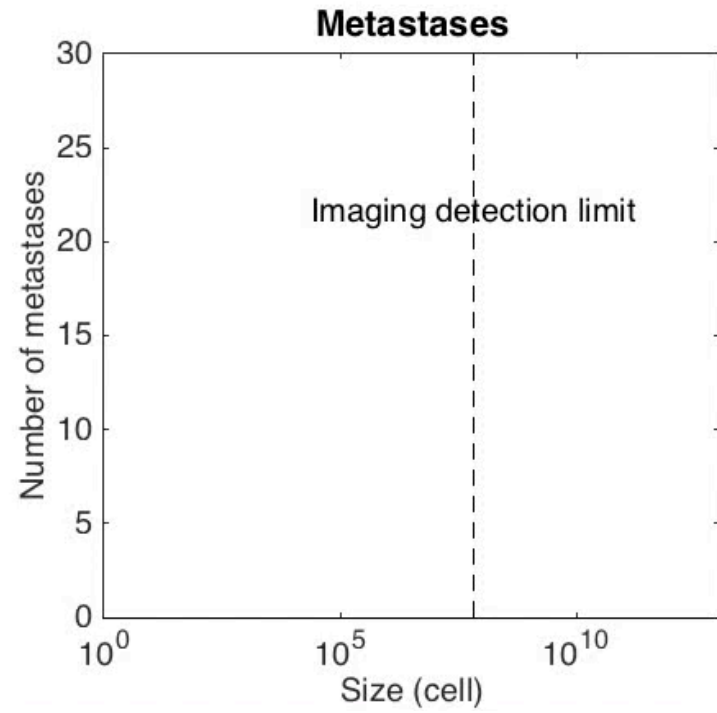
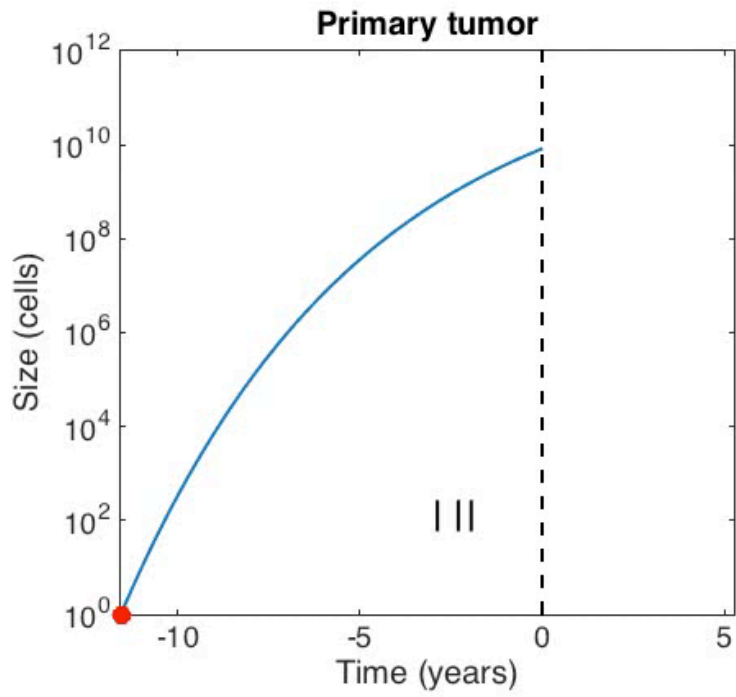
Parameter	Estimate	r.s.e. (%)	p-value
$\log \alpha_{pop}$	-8.883	10.151	
$\beta_{Ki67, \alpha}$	0.086	27.376	$2.59 \cdot 10^{-4}$
$\beta_{HER2, \alpha}$	0.029	42.833	0.020
$\beta_{CD44, \alpha}$	0.011	60.816	0.1
$\beta_{TRIO, \alpha}$	0.016	58.119	0.085
$\log \mu_{pop}$	-26.342	3.696	
$\beta_{EGFR, \mu}$	0.039	47.527	0.035
σ	0.606	23.104	
ω_{α}	2.062	22.715	
ω_{μ}	3.563	16.759	

c-index = 0.62 (10-folds cross-validation)

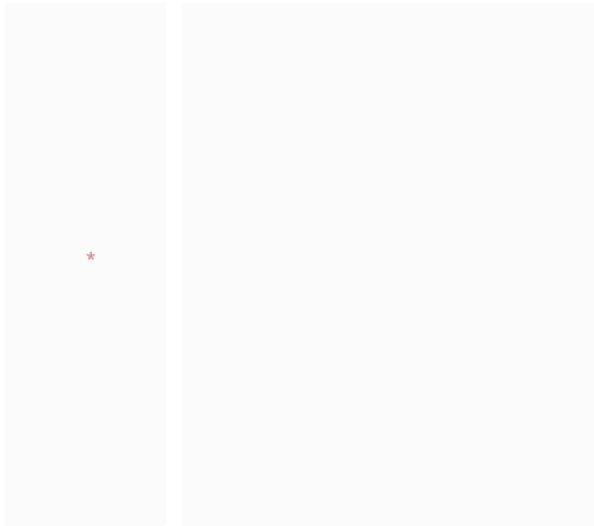


Patient ID	Tumor size (mm)	Ki67	HER2	CD44	TRIO	EGFR	Observed TTR (cens)	Predicted TTR	Prediction error (days)
47	20	32	100	0	0	50	739 (1)	447	292
255	25	1	60	90	60	0	1812 (1)	1609	203
143	18	60	0	50	0	0	2798 (1)	434	2364
12	10	20	0	23	0	0	5970 (0)	$+\infty$	-

$t = -11.6$ years

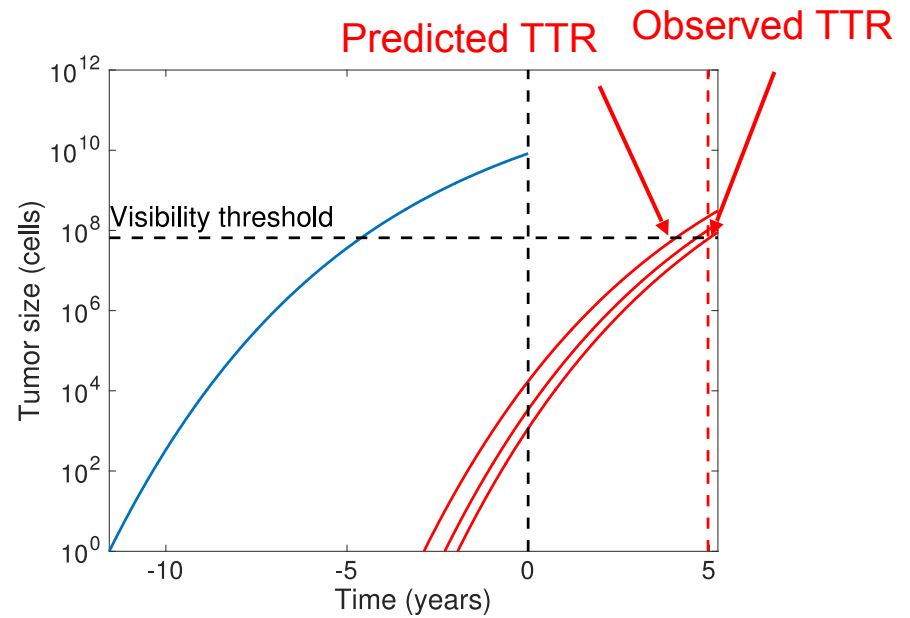


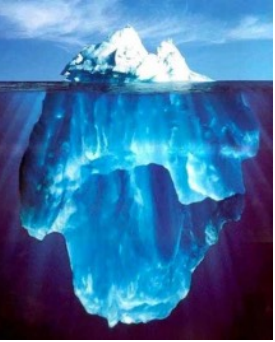
$t = -141$ months
— 10 mm



Primary tumor

Metastases



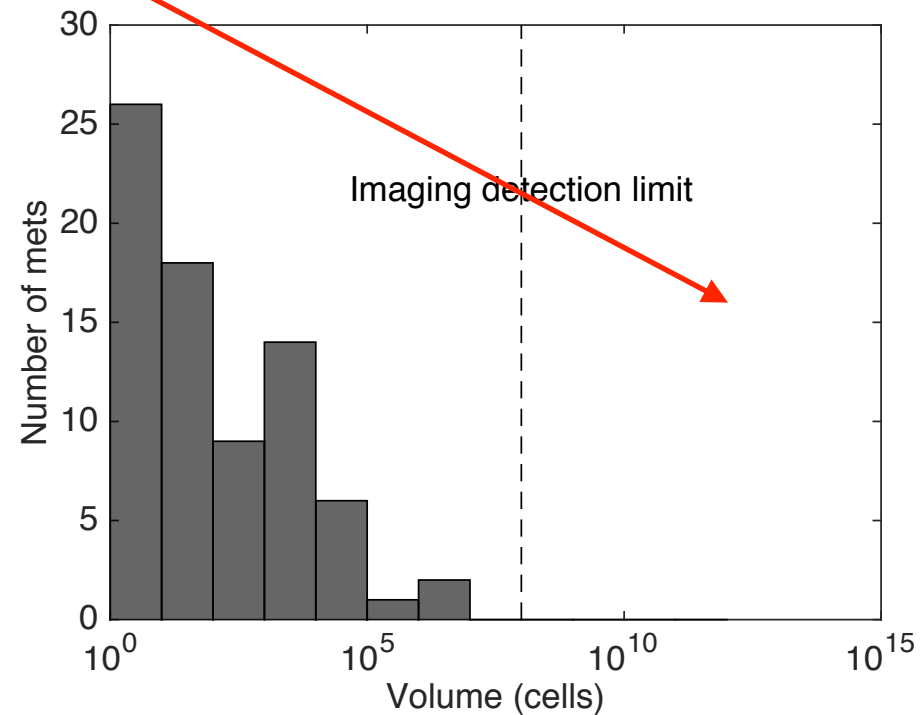
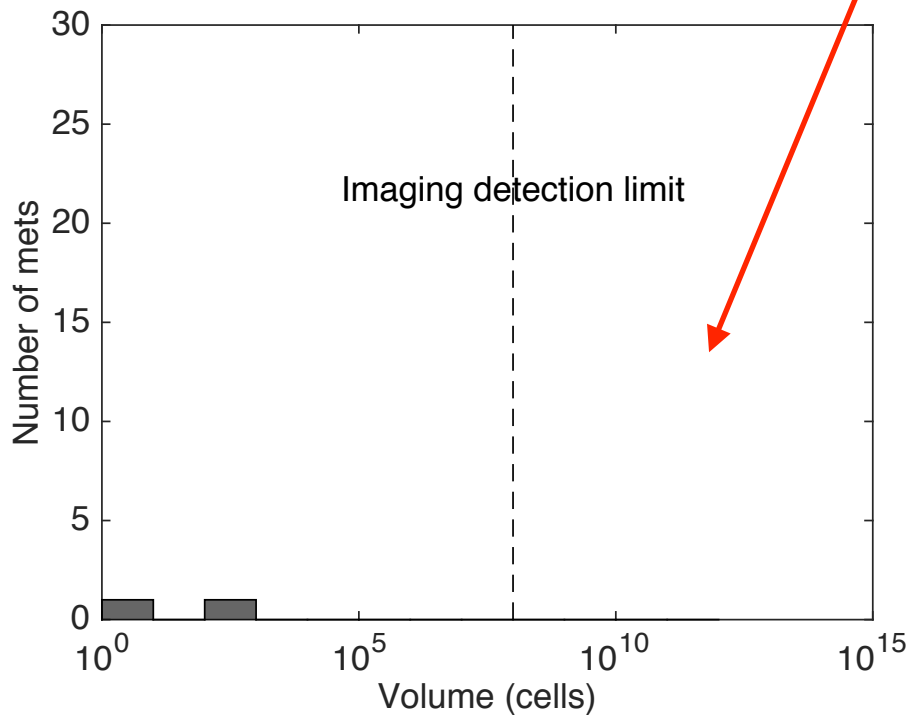


Diagnosis personalization

Virtual patient with
median μ

Virtual patient with
large μ (90th prct)

Nothing visible



Breast cancer patient with primary tumor of 4.32 cm

Chemotherapy personalization

Toward taking into account inter-individual variability

- 10 virtual patients with breast cancer detected at stage **T1N0M0**. Size of the tumor at detection: 1 gram.
- Chemotherapy : 6 cycles of 21 days (75mg of DTX and 100mg d'EPI) *Viens & al., J. Clin. Onc. 2001*
- Number of visible metastases ($> 10^8$ cel.) 5 years after the end of the treatment

Adapt the number of cycles to each patient

μ	Protocole de Viens		
	6 cycles 126 days	9 cycles 189 days	12 cycles 252 days
1.3×10^{-7}	1	0	*
2.7×10^{-7}	2	1	0
4.0×10^{-7}	3	2	1
6.1×10^{-7}	5	4	3

Acknowledgements

Biology

- Preclinical data of ortho-surgical animal models of metastases

*J. Ebos *A. Tracz
*M. Matri



Roswell Park Cancer Institute, Buffalo, NY, USA

- Beva + cytotoxics study



Dr. J. Ciccolini

Clinic

- Brain metastasis from lung tumors

*F. Chomy **Bergonié Institute, Bordeaux, FR**



*F. Barlesi

*X. Muracciole

AP-HM, Marseille, FR

Modeling

*C. Nicolò



*D. Barbolosi



That's all Folks!

Thanks for listening!

2 Combination bevacizumab - chemotherapy