# Bayesian Monitoring of Lapatinib (L) plus Trastuzumab (T) Treatment of Her2 Positive Metastatic Colorectal Cancer (mCRC): An Observational Cohort Study

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#### **BACKGROUND**

HER2 positivity is found in 3-5% of mCRC. The phase II HERACLES-A trial showed that dual anti-HER2 therapy with T + L has 30% response rate (RR) in HER2+ RAS wt mCRC after failure of standard care<sup>1,2</sup>. These data led to inclusion of T+ L among recommended treatments by NCCN and other guidelines, but still lack confirmation by large trials comparing treatments. The latter, especially

for this uncommon subset of mCRC, would require efforts unbearable in the setting of independent clinical research. We designed this study to confirm HERACLES-A data of efficacy through a Bayesian approach allowing to monitor longitudinally efficacy of T + L in the practice setting. Here we report the results of the first pre-planned interim-analysis.

### STATISTICAL DESIGN AND END POINTS

We simultaneously monitor efficacy and toxicity using the Bayesian optimal phase 2 (BOP2) design 3. Specifically, let n denote the interim sample size, and N denote the maximum sample size. Let Y<sub>eff</sub> and Y<sub>tox</sub> denote the binary efficacy and toxicity endpoints, with  $Y_{eff}=1$  and  $Y_{tox}=1$  indicating that experience efficacy and toxicity, respectively. We assume that the joint distribution of (Y<sub>eff</sub>, Y<sub>tox</sub>) follows a multinomial distribution with four elementary outcomes:  $(Y_{eff}, Y_{tox}) = (1, 1), (1, 0), (0, 1)$ 1) and (0, 0). Let  $p=(P_{11}, P_{10}, P_{01}, P_{00})$  denote the probabilities of observing the four outcomes, and let  $p_{eff}=Pr(Y_{eff}=1)$ ,  $p_{tox}=Pr(Y_{tox}=1)$  and  $p_{efftox}=Pr(Y_{eff}=1)$ Y<sub>tox</sub>=1). When efficacy and toxicity endpoints are monitored separately, the joint distribution reduced to marginal distribution of efficacy and marginal distribution of toxicity, respectively.

The treatment is deemed unacceptable if p<sub>eff</sub> ≤0.1 or  $p_{tox} > 0.3$ . Thus, we will stop enrolling patients and claim that the treatment is unacceptable if

 $Pr(p_{eff} > 0.1|data) < \lambda(n/N)^{\alpha}$ ,

 $Pr(p_{tox} \le 0.3|data) < \lambda(n/N)^{(\alpha/3)}$ 

where  $\lambda$ =0.5 and  $\alpha$ =0.51 are design parameters optimized to maximize the study power, i.e. probability of correctly concluding an efficacious and safe treatment as acceptable when  $p_{eff} = 0.35$ ,  $p_{tox} = 0.2$  and  $p_{efftox} = 0.07$ , while controlling that the probability of incorrectly claiming an inefficacious and toxic treatment, i.e., type I error, with  $p_{eff} = 0.1$  $p_{tox} = 0.3$  and  $p_{efftox} = 0.045$ , to 10.1%. Note that in the safety stopping rule, the original publication of the design used the probability cutoff  $\lambda(n/N)^{\alpha}$ , here the attenuation factor 3 is added (i.e.,  $\alpha/3$ ) to obtain stricter interim stopping boundaries to enhance

This optimization is performed assuming a vague Dirichlet prior *Dir(0.045,0.055,0.255,0.645)* for p. The prior is chosen such that it corresponds to a prior effective sample size of 1 patient, and the prior estimates of p<sub>eff</sub> and p<sub>tox</sub> match the values specified when the treatment is unacceptable. The above decision rule leads to the following optimal stopping

**Table 1: Optimized stopping boundaries** 

-	• • •	
# patients treated	Stop if # response ≤	OR # toxicity
10	0	4
20	1	7
30	2	10
40	4	13

The go/no-go criteria in Table 1 are non-binding.

Below are the operating characteristics of the design based on 10000 simulations using the BOP2 web application (BOP2 V1.4.6.0)<sup>3</sup>.

#### **Table 2: Operating characteristics**

			Pr(Eff	Early	Claim	Average
			&	Stopping	Acceptable	Sample
Scenario	Pr(Eff)	Pr(Tox)	Tox)	(%)	(%)	Size
1	0.10	0.3	0.045	82.03	10.09	18.4
2	0.35	0.2	0.070	18.37	80.75	35.2

# Study objectives **Primary**

• To evaluate ORR and safety of Her2 positive mCRC after Trastuzumab + Lapatinib

#### Secondary

 Progression free-survival and overall survival for this patient population

Patient's data and statistical analysis will be reported according to most updated guidelines<sup>4</sup>.

This observational study was approved by our institutional EC.

#### **METHODS AND PATIENTS**

Patients inclusion criteria were the same from Heracles trial<sup>1</sup>. In addition, since this is on observational study, the patients should be already candidate to off-label T + L treatment at our institution according to Italian and hospital rules, after acknowledgment from our EC. The Division of Sex Oncology and Fondazione Oncologia Niguarda (FON) covered the costs of treatments along with AIFA funding, according to law 326/2003 (5% AIFA National Fund).

Efficacy evaluation. Patients not evaluable for ORR will be replaced. The Efficacy Evaluable (EE) population is defined as all treated patients, with no major deviations from the eligibility criteria affecting efficacy evaluation, for whom the tumor response could be evaluated at least once while on treatment. These patients should have received at least 2 cycles after treatment starts, unless disease

# RESULTS (I)

From May 2019 to Jan 2021, we collected data of HER2+ mCRC patients treated with T + L according to HERACLES trial inclusion criteria. 11 patients were followed for RR and adverse events (AE). On May 1st 2021, at the first interim analysis, 2/10 evaluable patients had PR according to RECIST criteria, with an updated probability of response of 25.8% (Figure 2). Figure 1 shows an example of response in a patient with a large metastatic liver lesion. No G3 AE drug-related were reported meeting the safety boundaries of the observational protocol. All AEs are detailed in Table 4 (right).

Prior cetuximab or panitumumab Prior anti-HER2 therapy Subsequent treatment Surgery Table 4: AE related to study drug Grades 1-2 n(%) Gastrointestinal 11(100) Vomiting 1 (9) **Dermatological** Folliculitis 2 (18)

2.3 (1-4)

progression occurs within this period. Patients

characteristics are reported in Table 3.

**Table 3: Patients characteristics.** 

Median age in years (range)

**HER2** expression by IHC score

Trasversum, left colon and rectum

Median N of prior lines (range)

Patients with > 3 prior lines

Sideness of primary tumor

**Prior treatment** 

2 (18) Mucositis Metabolic and nutritional disorders Anorexia Astenia 1 (9) Bilirubin increased 2 (18)

Figure 2 – Tumor shrinkage in a liver metastasis of a patient with mCRC Her2 positive, after 2 months of T +L treatment (PR





# **RESULTS (II)**

Figure 1: Updated probability of response to T + L in Her2 positive mCRC, after first interim analysis

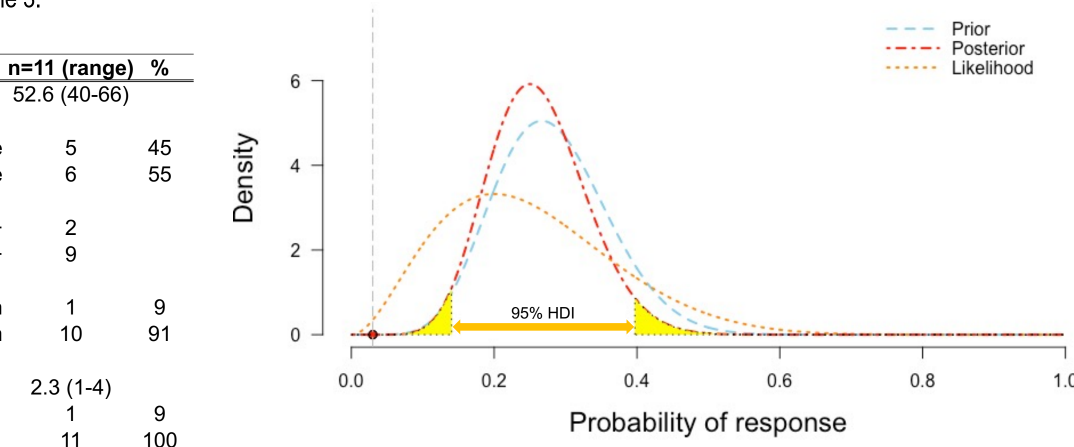


Figure 1 (left) shows the prior probabilities of response (blue) from published data<sup>4</sup>, the likelihood of response observed in the present study and the update posterior probability density (red) along with the 95% high density interval (HDI, 13-39%). Response to T + L in Her2 pos mCRC has a strong evidence to be better than the available choices in 3<sup>rd</sup> line 1.0 (3% response rate H0, red point), with a  $BF_{10}$  of 7.401.

On the right, the swimmer plot shows the response rate (color code) and progression free survival to T+L treatment (ranging from 2 to 21 months) for each individual patient. One patient (#007) underwent to liver metastasectomy right after response to the study treatment and is alive and progression free at the interim analysis

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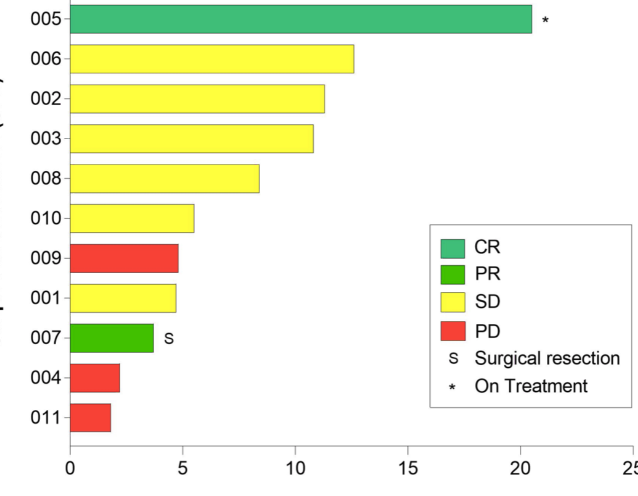


Figure 3: Swimmer plot showing RR and PFS

#### **CONCLUSIONS**

effective. A Bayesian approach, allowing to monitor HER2+ mCRC. results accounting for previously available data, can

At the first planned interim analysis, treatment with T + L support the process of approval by regulatory authorities for HER2+ mCRC patients was confirmed to be safe and for treatments targeted to uncommon subsets such as

Progression-free survival (months)

## **REFERENCES**

- Sartore-Bianchi A, Trusolino L, Martino C, et al: Dual-targeted therapy with 3. Zhou, Heng, J. Jack Lee, e Ying Yuan. 2017. «BOP2: Bayesian Optimal Design trastuzumab and lapatinib in treatment-refractory, KRAS codon 12/13 wild-type, HER2-positive metastatic colorectal cancer (HERACLES): a proof-of-concept, multicentre, open-label, phase 2 trial. Lancet Oncol 17:738-746, 2016. https://doi.org/10.1016/S1470-2045(16)00150-9
- 2. Tosi, Federica, Andrea Sartore-Bianchi, Sara Lonardi, Alessio Amatu, Francesco Leone, Silvia Ghezzi, Cosimo Martino, et al. 2020. «Long-Term Clinical Outcome of Trastuzumab and Lapatinib for HER2-Positive Metastatic Colorectal Cancer». Clinical Colorectal Cancer 19 (4): 256-262.e2. https://doi.org/10.1016/j.clcc.2020.06.009.
- for Phase II Clinical Trials with Simple and Complex Endpoints». Statistics in Medicine 36 (21): 3302–14. https://doi.org/10.1002/sim.7338
- 4. Schoot, Rens van de, Sarah Depaoli, Ruth King, Bianca Kramer, Kaspar Märtens, Mahlet G. Tadesse, Marina Vannucci, et al. 2021. «Bayesian Statistics and Modelling». Nature Reviews Methods Primers 1 (1): 1–26. https://doi.org/10.1038/s43586-020-00001-2





