

Post-resection/ablation chemotherapy in patients with metastatic colorectal cancer (FIRE-9 - PORT/ AIO KRK0418)

Prospective, randomized, open, multicenter Phase III trial to investigate the efficacy of active post-resection/ablation therapy in patients with metastatic colorectal cancer

Investigational medicinal products	5-FU, folinic acid, irinotecan, oxaliplatin
EudraCT no.	2020-006144-18
Protocol code	FIRE-9 - PORT
AIO trial-no.	AIO-KRK 0418
Short title	mFOLFOXIRI/FOLFOX versus follow- up surveillance after definite treatment of colorectal cancer metastases
Sponsor	Charité Universitätsmedizin Berlin Charitéplatz1 10117 Berlin
Sponsor representative	Prof. Dr. med. Dominik Modest
Version Draft 0.5	14-December-2020
Amendment No.	

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Responsibilities

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Approval of the Protocol

Trial title: Post-resection/ablation chemotherapy in patients with metastatic colorectal cancer (FIRE-9 - PORT/ AIO KRK0418)

Prospective, randomized, open, multicenter Phase III trial to investigate the efficacy of active post-resection/ablation therapy in patients with metastatic colorectal cancer

EudraCT no.: 2020-006144-18

I have approved the protocol version 0.1 dated 14-December-2020 and confirm that the clinical trial will be conducted in accordance with this protocol, the Declaration of Helsinki, the International Conference on Harmonization (ICH) for Good Clinical Practice (GCP) guidelines and the applicable regulatory requirements.

I furthermore confirm that the investigators and institutions involved in the clinical trial are to permit clinical trial-related monitoring, audits and regulatory inspections, including provision of direct access to source data and documents.

Representative of the Sponsor

Name of sponsor representative

Date

Signature

Approval of the Protocol

Trial title: Prospective, randomized, open, multicenter Phase II trial to investigate the efficacy of trifluridine/tipiracil plus panitumumab versus trifluridine/tipiracil plus bevacizumab as first-line treatment of metastatic colorectal cancer

EudraCT no.: 2020-006144-18

I have approved the protocol version 0.5 dated 22-September-2020 and confirm that the clinical trial will be conducted in accordance with this protocol, the Declaration of Helsinki, the International Conference on Harmonization (ICH) for Good Clinical Practice (GCP) guidelines and the applicable regulatory requirements.

Coordinating Investigator

Name of coordinating investigator

Date

Signature

Approval of the Protocol

Trial title: Post-resection/ablation chemotherapy in patients with metastatic colorectal cancer (FIRE-9 - PORT/ AIO KRK0418)

Prospective, randomized, open, multicenter Phase III trial to investigate the efficacy of active post-resection/ablation therapy in patients with metastatic colorectal cancer

EudraCT no.: 2020-006144-18

I have approved the protocol version 0.1 dated 14-December-2020 and confirm that the clinical trial will be conducted in accordance with this protocol, the Declaration of Helsinki, the International Conference on Harmonization (ICH) for Good Clinical Practice (GCP) guidelines and the applicable regulatory requirements.

Biometrician

Name of the biometrician

Date

Signature

Statement of Compliance

Trial title: Post-resection/ablation chemotherapy in patients with metastatic colorectal cancer (FIRE-9 - PORT/ AIO KRK0418)

Prospective, randomized, open, multicenter Phase III trial to investigate the efficacy of active post-resection/ablation therapy in patients with metastatic colorectal cancer

EudraCT no.: 2020-006144-18

I have read and understood this protocol and agree to conduct the clinical trial in accordance with this protocol, the Declaration of Helsinki, the International Conference on Harmonization (ICH) for Good Clinical Practice (GCP) guidelines and the applicable regulatory requirements.

I agree to ensure that the confidential information contained in this document will not be used for any purpose other than the evaluation or conduct of the clinical trial without the prior written consent of the sponsor of the clinical trial.

Investigator

Signature

Date

Investigator

Investigator's Institution

Title	Post-resection/ablation chemotherapy in patients with metastatic colorectal cancer (FIRE-9 - PORT/ AIO KRK0418). Prospective, randomized, open, multicenter Phase III trial to investigate the efficacy of active post-resection/ablation therapy in patients with metastatic colorectal cancer
Short title	mFOLFOXIRI/FOLFOX versus follow-up surveillance after definite treatment of colorectal cancer metastases
Sponsor	Charité Universitätsmedizin Berlin Charitéplatz1 10117 Berlin
Coordinating investigator	Prof. Dr. med. Dominik Modest Charité - Universitätsmedizin Berlin Medizinische Klinik mit Schwerpunkt Hämatologie, Onkologie und Tumorimmunologie am Campus Virchow Klinikum (CVK) Augustenburger Platz 1 13353 Berlin
Protocol code	FIRE-9 – PORT
AIO trial-no.	AIO KRK 0418
EudraCT no	2020-006144-18
version and version date	Draft 0.1 of 14-December-2020
Investigational medicinal product(s)	5-FU, folinic acid, irinotecan, oxaliplatin
Clinical trial phase	Phase III
Number of patients	507 patients
Number of trial sites	About 80 trial sites
Indication studied	Post-resection/ablation chemotherapy in patients with metastatic colorectal cancer
Background and rationale	In Germany, colorectal cancer has a prevalence of 65-80/100.000 and a current 5- year mortality of appr. 50% (Robert-Koch-Institut: Krebsdaten 2015; www.krebsdaten.de). In Western Europe, the burden of colorectal cancer is reported to be 211 (female) and 298 (male) diability adjusted life years (DALYs) on a population of 100.000 2. Given the available screening-programs, no severe socioeconomic impact within the incidence appears present. Life-style attitudes however may affect individual risk.
	Patients with metastases from colorectal cancer (appr. 40-50% of all patients develop metastases) benefit from the resection or ablation of metastates, although relapse occurs in the majority (appr. 70-80%) of these patients 3-7. Clearly, a reduction in relapse rates would improve the long-term outcome of these patients.
	Unfortunately, additive/adjuvant therapy after local treatment of metastases is not established by phase III trials. Accordingly, no standard of care treatment to improve the relapse rates is available and the current S3-guideline for colorectal cancer does not recommend additive chemotherapy due to insufficient evidence on its benefit (http://www.awmf.org/leitlinien/detail/II/021-007OL.html), explicitly. The present clinical trial aims to generate evidence that additive therapy after resection or ablation of metastases may improve PFS and OS in patients with colorectal cancer. This is of

Synopsis

Objectives	 specific importance since both improvements in in localized, but also systemic therapies have resulted in increasing numbers of mCRC patients undergoing resection and/or ablation of metastases4,8-13. Optimal oncological management after removal of metastases is unclear. The result of this trial may be therefore be practice-changing. To support the purely clinical information a supporting translational study will help to identify subgroups (if present) of patients that benefit/ or not from systemic therapy after removal of metastases. Primary objective To compare the efficacy of active additive chemotherapy after definitive
	 treatment of metastases to structured follow-up only <u>Secondary objectives</u> To compare efficacy, safety and patient reported quality of life (QoL) of active treatment to structured follow-up only
	Other exploratory objective Further anti-tumor treatment after discontinuation of study treatment
	<u>Translational research objectives</u> Identification and characterization of patient subgroups with greatest or lowest benefit from treatment including efficacy and toxicity using tumor specimen and blood-based biomarker candidates.
Endpoints	 Primary endpoint Progression-free survival (PFS) time at the 24 months follow-up defined as time from randomization to death or evidence of disease as defined by RECIST 1.1 criteria (whatever occurs first) Secondary endpoints
	 Efficacy PFS in patients with/without prior systemic therapy PFS in patients with R1 vs R0 resected lesions as well as ablated vs purely resected lesions overall survival safety treatments (including efficacy) beyond study participation
	 local control of lesions according to ablative technique (surgery vs. ablation vs. radiation). Local PFS?
	Quality of lifeQoL as assessed with the QoL questionnaire EQ-5D-5L
	 Safety Type, incidence, severity, and causal relationship to active chemotherapy of non-serious adverse events and serious adverse events (severity evaluated according to CTCAE version 5.0)
	Other exploratory endpoints Translational analyses including evaluation of tumor specimen of primary and/or metastatic tissue as well as blood samples at different time points
	 PFS and OS according to circulating tumor DNA at baseline (ctDNA positive vs negative), outcome in molecular subgroups,



	6. ECOG performance status 0-2
	7. Adequate bone marrow, hepatic and renal organ function, defined by the
	following laboratory test results:
	 Absolute neutrophil count ≥ 1.5 x 10⁹/L (1500/µL)
	 Hemoglobin ≥ 80 g/L (8 g/dL)
	 Platelet count ≥ 100 x10⁹/L (75,000/µL) without transfusion
	 Total serum bilirubin of ≤ 1.5 x upper limit of normal (ULN)
	 Aspartate aminotransferase (AST/GOT) and alanine aminotransferase (ALT/GPT) ≤ 3.0 × ULN
	 Calculated glomerular filtration rate (GFR) according to Cockcroft –Gault formula or according to MDRD ≥ 50 mL/min or serum creatinine ≤ 1.5 x ULN
	 Patients without anticoagulation need to present with an INR <1.5 x ULN and PTT <1.5 x ULN.
	9. Proficient fluorouracil metabolism as defined:
	 a) Prior treatment with 5-FU or capecitabine without unusal toxicity or
	b) Normal DPYD deficiency test according to the standard of the study centre
	10. For females of childbearing potential (FCBP): negative pregnancy test within
	14 days before randomisation and agreement to remain abstinent (refrain from
	heterosexual intercourse) or use contraceptive methods with a failure rate of
	<1% per year during the treatment period and for at least 6 months after the last dose of study treatment
	A woman is considered to be of childbearing potential if she is post-
	menarcheal, has not reached a postmenopausal state (\geq 12 continuous
	months of amenorrhea with no identified cause other than menopause), and
	has not undergone surgical sterilization (removal of ovaries and/or uterus).
	Examples of contraceptive methods with a failure rate of <1% per year include
	bilateral tubal ligation, male partner's sterilization, hormonal contraceptives that
	inhibit ovulation, hormone-releasing intrauterine devices, and copper
	Intrauterine devices.
	abstinent or use a condom plus an additional contracentive method that
	together result in a failure rate of $<1\%$ per year during the treatment period and
	for 6 months after the last dose of study treatment. Men must refrain from
	donating sperm during this same period.
	With pregnant female partners, men must remain abstinent or use a condom
	during the treatment period and for 6 months after the last dose of study
	medication to avoid exposing the embryo.
Exclusion criteria	1. Treatment of metastases greater than 3cm with radio-frequency/microwave ablation within 24 months prior to study entry if applicable
	2. Treatment of lesions greater than 5cm with radiation (stereotactic/ brachytherapy) within 24 months prior to study entry if applicable
	 Previous chemotherapy for metastatic or localized disease with >6 cycles of FOLFOX (or FOLFOXIRI) or >4 cycles of CAPOX/XELOX
	4. New York Heart Association Class III or greater heart failure by clinical
	judgement
	5. Myocardial infarction within 6 months prior to randomisation; percutaneous
	transluminal coronary angioplasty (PTCA) with or without stenting within 6
	months prior to randomization
	6. Unstable angina pectoris
	7. Unstable cardiac arrnythmia > grade 2 NCI CI CAE despite anti-arrhythmic therapy
	8 Ongoing toxicities > grade 2 NCL CTCAF in particular peripheral neuropathy
	o. Ongoing toxiolities > grade 2 nor of OAE, in particular peripheral neuropatiny

	9. Active uncontrolled infection
	 Severe chronic non-healing wounds, ulcerous lesions or untreated bone fracture.
	11. Known hypersensitivity to 5-FU, folinic acid, irinotecan or oxaliplatin
	12. Major surgical procedure, open biopsy, or significant traumatic injury within 21 days prior to randomisation, or abdominal surgery, abdominal interventions or significant abdominal traumatic injury within 21 days prior to randomisation or anticipation of need for major surgical procedure during the course of the study
	or non-recovery from side effects of any such procedure
	13. Any other disease, metabolic dysfunction, physical examination finding, or clinical laboratory finding that contraindicates the use of an investigational drug, may affect the interpretation of the results, or may render the patient at high risk from treatment complications.
	14 Medical history of other malignant disease than mCPC with the following
	exceptions:
	 patients who have been disease-free for at least three years before randomisation
	 patients with adequately treated and completely resected basal cell or squamous cell skin cancer, in situ cervical, breast or prostate cancer, stage I uterine cancer
	 patients with any treated or untreated malignant disease that is associated with a 5 year survival prognosis of ≥90% and does not require active therapy
	15. Known alcohol or drug abuse
	16. Pregnant or breastfeeding females
	17. Participation in a clinical trial or experimental drug treatment within 28 days prior to potential inclusion in the clinical trial or within a period of 5 half-lives of the substances administered in a clinical trial or during an experimental drug treatment prior to potential inclusion in the clinical trial, depending on which period is longest, or simultaneous participation in another clinical trial while taking part in this clinical trial.
	 Patient committed to an institution by virtue of an order issued either by the iudicial or the administrative authorities
	19. Limited legal capacity
Treatment,	Arm A:
dosage and administration	 14-day cycle, choice of (by stratification) intravenous therapy 1. mFOLFOXIRI: (d1: 2.4g 5-FU in 46 hours, 400mg/qm leucovorin, 85mg/qm oxaliplatin, 150mg/qm irinotecan)
	 mFOLFOX: (d1: 2.4g 5-FU in 46 hours, 400mg/qm leucovorin, 85mg/qm oxaliplatin)
	<i>De-escalations and dose modifications are allowed per institutional standard as investigators decision.</i>
	Arm A and B: Radiologic re-assessment with computed tomography is scheduled 3 months (6 cycles of therapy or 3 months observation) and 6 months (12 cycles of therapy or 6 months of observation) after randomisation
	Structured follow-up for up to 60 months after randomization should be maintained. It is recommended to offer CT scans of thorax/abdomen and/or MRI scans every 3 months within the 2 years after randomisation (6 months =2 controls are part of the study. After the first two relapse-free years, intervals are stretched to 6 months in the third and following years after study participation.

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	The treatment is continued in Arm A for a maximum of 12 cycles or until progression according to RECIST 1.1 criteria or unacceptable toxicity whatever comes first.
Translational research	Collection of primary and metastastic tumor tissue and 3 blood sample time points (baseline, 3months, 6 months)
Statistical considerations	The primary endpoint will be PFS (progression-free survival, defined as progression/relapse or death from any cause) at 30 months after randomization. Our assumptions are based on the only available analysis of two pooled, early-stopped trials in the adjuvant/additive setting using fluoropyrimidine monotherapies18 that reported a hazard ratio for PFS in favor of active treatment of 0.76 (the originally reported hazard ratio was reported reversed as 1.32) that translated into a similar effect for the endpoint overall survival. This again support the validity of the primary endpoint PFS in this application. We hypothesize a slightly larger effect for PFS in favor of active therapy due to the 2-3 drug regimens resulting in an estimated hazard ratio of 0.70. For the PORT trial18 for the control arm (surgery alone) with structured follow-up, a progression/relapse/death-free rate of 40% at time point 24 months was observed translating into a 60% progression/relapse/death-rate at this time (according to the reported control arm 18). With a hazard ratio 0.70 (= $\Box \Box \Box C C = 0.0267/0.0382$) favoring active treatment, the hypothesized relapse rate at 24 months in the intervention arm is assumed to be: 47%. With a power of 80%, a 2-sided alpha of 0,05, a total of 276 events are need to be observed in order to detect a difference in progression-free survival of a hazard ratio of 0.70-favoring active treatment vs. observation (Schoenfeld formula). Assuming an accrual time of 48 months and a follow-up time of 24 months, a drop-out/censoring rate of 40% after 24 months fire randomization, a total of 480 patients (320/160 in the respective arms, rounded to receive integers and maintain the allocation ratio) is expected to yield the required number of events if the accrual rate is constant. The computation was done using the software R Version 3.5.1 and the package Rpact. We account for additional 5% of patients that directly leave the study after randomization and never received the study medication. Thus a total of 507 patie
Duration and end of trial	ca. 4-5 years recruitment and up to 5 years follow-up per patient
GCP statement	The clinical trial will be conducted in accordance with the ethical principles that have their origin in the Declaration of Helsinki and that are consistent with the International Conference on Harmonization (ICH) for Good Clinical Practice (GCP) and the applicable regulatory requirements.

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