# Phase I study of S 95005 in combination with oxaliplatin in metastatic colorectal cancer

Submission date	<b>Recruitment status</b> No longer recruiting	[X] Prospectively registered		
02/02/2016		☐ Protocol		
Registration date	Overall study status Completed	Statistical analysis plan		
02/03/2016		[X] Results		
Last Edited	Condition category	Individual participant data		
23/06/2021	Cancer			

## Plain English summary of protocol

Not provided at time of registration and not expected to be available in the future

# Contact information

# Type(s)

Scientific

### Contact name

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### Contact details

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# Type(s)

Public

### Contact name

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### Contact details

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# Additional identifiers

# Clinical Trials Information System (CTIS)

2015-004894-34

# ClinicalTrials.gov (NCT)

NCT02848443

### Protocol serial number

CL1-95005-001

# Study information

### Scientific Title

Phase I dose-escalation of S 95005 (TAS-102) in combination with oxaliplatin in metastatic colorectal cancer

### **Study objectives**

To assess the safety and tolerability and to determine the recommended phase 2 dose of S 95005 given in combination with oxaliplatin in patients with metastatic colorectal cancer.

### Ethics approval required

Old ethics approval format

# Ethics approval(s)

Comité Ético de Investigación Clínica del Hospital Universitari Vall d'Hebron, 04/03/2016

# Study design

Multicentre open-label non-randomised non-comparative study

# Primary study design

Interventional

# Study type(s)

Treatment

# Health condition(s) or problem(s) studied

Metastatic colorectal cancer

### **Interventions**

This is a one-arm study, which will be conducted in 2 parts:

- 1. A dose-escalation part to determine the maximum tolerated dose (MTD) of S 95005 in combination with oxaliplatin: a minimum of 3 patients will be enrolled at the initial dose level of 25 mg/m<sup>2</sup> of S95005 in combination with 85 mg/m<sup>2</sup> of oxalipatin. Patients will be included by groups of 3.
- 2. An expansion part in patients treated at the recommended dose defined in the dose escalation part of this study to evaluate the safety, PK, and preliminary efficacy of S 95005 in combination with oxaliplatin and either bevacizumab or nivolumab.

The treatments will be given until unacceptable toxicity according to the investigator, disease

progression or patient withdrawal. The follow-up will last up to 6 months after the end of the participation in the study.

S95005: film-coated tablets containing 15mg of trifluridine and 7.065mg of tipiracil hydrochloride, or 20mg of trifluridine and 9.42mg of tipiracil hydrochloride, given orally at the dose of 25 or 30 or 35 mg/m2/dose

Oxaliplatin: concentrate for solution for infusion containing 5mg/ml of oxaliplatin, administered intravenously at the dose of 65 to 85 mg/m2

Bevacizumab: concentrate for solution for infusion containing 25mg/ml of bevacizumab, administered intravenously at the dose of 5 mg/kg

### Added 05/04/2017:

Nivolumab: concentrate for solution for infusion containing 10 mg/ml of nivolumab, administered intravenously at the dose of 3 mg/kg.

### Intervention Type

Drug

### Phase

Phase I

### Drug/device/biological/vaccine name(s)

1. S95005 (trifluridine/tipiracil hydrochloride) 2. Oxaliplatin 3. Bevacizumab 4. Nivolumab

# Primary outcome(s)

- 1. Maximum tolerated dose (MTD) and dose limiting toxicity (DLT) of S95005 when given in combination with oxaliplatin, during the first two cycles in the dose-escalation part
- 2. Safety tolerance profile of S 95005 given in combination with oxaliplatin, at each visit, from the informed consent signature to the withdrawal visit, assessed by: adverse events, physical examinations and ECOG performance status, laboratory examinations (haematology, biochemistry and urinalysis), vital signs, ECG and body weight

### Key secondary outcome(s))

Secondary outcome measures as of 05/04/2017:

- 1. Main pharmacokinetic parameters of S 95005 and its main metabolites, and oxaliplatin, from day 1 of cycle 1 to day 5 of cycle 2
- 2. Antitumor activity (objective response rate, duration of response, Progression-free survival and Overall survival) assessed by RECIST (Response Evaluation Criteria in Solid Tumors) and CEA (Carcinoembryonic Antigen), from the informed consent signature to the withdrawal visit
- 3. Safety tolerance profile of S 95005 in combination with oxaliplatin and either bevacizumab or nivolumab assessed by: adverse events, physical examinations and performance status, laboratory examinations (haematology, biochemistry and urinalysis), vital signs and body weight, from the informed consent signature to the withdrawal visit
- 4. PDL-1 expression, tumour-infiltrating CD8 T cell density: tumor biopsy at baseline and at the end of cycle 4
- 5. Exploratory endpoints: Proteomic and genomic biomarkers using blood samples, after consent signature from day 1 of cycle 1 to the withdrawal visit (all patients), and tumour biopsies (for patients receiving nivolumab, at baseline and at the end of cycle 4)

Original secondary outcome measures:

- 1. Main pharmacokinetic parameters of S 95005 and its main metabolites, and oxaliplatin, from day 1 of cycle 1 to day 5 of cycle 2
- 2. Antitumor activity (objective response rate, duration of response, Progression-free survival and Overall survival) assessed by RECIST (Response Evaluation Criteria in Solid Tumors), from the informed consent signature to the withdrawal visit
- 3. Exploratory endpoints: Proteomic and genomic biomarkers using blood samples, after consent signature from day 1 of cycle 1 to the withdrawal visit

### Completion date

09/04/2020

# **Eligibility**

### Key inclusion criteria

Inclusion criteria as of 24/11/2016:

- 1. Age 18 years or older
- 2. Histologically confirmed metastatic colorectal cancer pretreated by at least one line of standard chemotherapy
- 3. Restaging scan within 28 days before the first study drug intake
- 4. During the dose-escalation part, patient must have at least one evaluable or measurable metastatic lesion; and during the expansion part, patient must have at least one measurable metastatic lesion
- 5. Life expectancy of more than 3 months
- 6. Performance status Eastern Cooperative Oncology Group (ECOG): 0-1
- 7. Adequate bone marrow, liver, and kidney function
- 8. For patient who will receive bevacizumab: coagulation parameters in normal limit or in therapeutic limit for patients treated with anticoagulant
- 9. Women of childbearing potential must have a negative pregnancy test. Female participants of childbearing potential and male participants with partners of childbearing potential must agree to use highly effective birth control method. Women and female partners using hormonal contraceptive must also use a barrier method.
- 10. Capacity to take oral tablet(s) without difficulty
- 11. Has provided written informed consent
- 12. Is willing and able to comply with scheduled visits and study procedures

Added 06/04/2017: For patients who will receive nivolumab: patients eligible for tumour biopsy and who agree to have two sequential biopsies during the study

### Original inclusion criteria:

- 1. Age 18 years or older
- 2. Histologically confirmed metastatic colorectal cancer pretreated by at least one line of standard chemotherapy and naïve to oxaliplatin in the metastatic setting
- 3. Restaging scan within 28 days before the first study drug intake
- 4. During the dose-escalation part, patient must have at least one evaluable or measurable metastatic lesion; and during the expansion part, patient must have at least one measurable metastatic lesion
- 5. Life expectancy of more than 3 months
- 6. Performance status Eastern Cooperative Oncology Group (ECOG): 0-1
- 7. Adequate bone marrow, liver, and kidney function
- 8. For patient who will receive bevacizumab: coagulation parameters in normal limit and

adequate proteinuria

- 9. Women of childbearing potential must have a negative pregnancy test Both males and females must agree to use effective birth control method
- 10. Capacity to take oral tablet(s) without difficulty
- 11. Has provided written informed consent
- 12. Is willing and able to comply with scheduled visits and study procedures

### Participant type(s)

Patient

### Healthy volunteers allowed

No

### Age group

Adult

### Lower age limit

18 years

### Sex

All

### Total final enrolment

78

## Key exclusion criteria

Exclusion criteria as of 05/04/2017:

- 1. Grade 2 or higher peripheral neuropathy
- 2. During expansion part, patients who had recurrence during or within 6 months of completion of the adjuvant chemotherapy with oxaliplatin
- 3. Patients with brain metastases or leptomeningeal metastasis
- 4. Other active malignancy within the last 3 years (except for basal cell carcinoma or a non-invasive/in situ cervical cancer)
- 5. Has had certain other recent treatment e.g. major surgery, field radiation, participation in another interventional study within the specified time frames prior to study drug administration 6. For patient who will receive bevacizumab: history of allergic reactions/hypersensitivity to bevacizumab to any components used in the formulation, to Chinese Hamster Ovary (CHO) cell
- products or other recombinant human or humanised antibodies.

  7. Grade 3 or higher hypersensitivity reaction to oxaliplatin, or grade 1.
- 7. Grade 3 or higher hypersensitivity reaction to oxaliplatin, or grade 1-2 hypersensitivity reaction to oxaliplatin not controlled with premedication
- 8. Patient previously treated by S 95005 or history of allergic reactions attributed to compounds of similar or biologic composition to S 95005 or any of its excipient, or has rare hereditary problems of galactose intolerance, the Lapp lactase deficiency or glucose-galactose malabsorption.
- 9. Certain serious illnesses or serious medical conditions
- 10. Any condition that, in the judgment of the Investigator, may affect the patient's ability to understand and sign the informed consent and fully comply with all study procedure
- 11. Pregnancy or breast feeding
- 12. For patients planned to receive nivolumab:
- 12.1. Patients with active autoimmune disease or history of clinically severe autoimmune disease.
- 12.2. Patients with a condition requiring systemic treatment with either corticosteroids (> 20 mg

daily prednisone equivalent) or other immunosuppressive medications within the specified time frames prior to first study drugs intake.

- 12.3. Prior treatment with anti-PD-1, anti-PD-L1, anti-programmed cell death ligand-2, anti-CD137, anti-OX-40, anti-CD40, anti-cytotoxic T lymphocyte-associated antigen-4 antibodies (CTLA-4), or any other immune checkpoint inhibitors.
- 12.4. Prior events of immune-mediated pneumonitis, immune-mediated colitis, immune-mediated hepatitis, immune-mediated endocrinopathies, immune-mediated nephritis and renal dysfunction, immune-mediated rash, immune-mediated encephalitis.
- 12.5. Allergic reactions/hypersensitivity to nivolumab or any components used in its formulation or previous severe hypersensitivity reaction to treatment with another monoclonal antibody. 12.6. Has a known history of active tuberculosis (Bacillus Tuberculosis).

### Exclusion criteria as of 24/11/2016:

- 1. Grade 2 or higher peripheral neuropathy
- 2. During expansion part, patients who had recurrence during or within 6 months of completion of the adjuvant chemotherapy with oxaliplatin
- 3. Patients with brain metastases or leptomeningeal metastasis
- 4. Other active malignancy within the last 3 years (except for basal cell carcinoma or a non-invasive/in situ cervical cancer)
- 5. Has had certain other recent treatment e.g. major surgery, field radiation, received investigational agent, within the specified time frames prior to study drug administration 6. History of allergic reactions/hypersensitivity to bevacizumab (for patient who will receive bevacizumab) or any components used in the formulation
- 7. Grade 3 or higher hypersensitivity reaction to oxaliplatin, or grade 1-2 hypersensitivity reaction to oxaliplatin not controlled with premedication
- 8. Patient previously treated by S 95005 or history of allergic reactions attributed to compounds of similar or biologic composition to S 95005 or any of its excipient, or has rare hereditary problems of galactose intolerance, the Lapp lactase deficiency or glucose-galactose malabsorption.
- 9. Certain serious illnesses or serious medical conditions
- 10. Any condition that, in the judgment of the Investigator, may affect the patient's ability to understand and sign the informed consent and fully comply with all study procedure 11. Pregnancy or breast feeding

### Original exclusion criteria:

- 1. Grade 2 or higher peripheral neuropathy
- 2. Patients who had recurrence during or within 6 months of completion of the adjuvant chemotherapy with oxaliplatin
- 3. Patients with brain metastases or leptomeningeal metastasis
- 4. Other active malignancy within the last 3 years (except for basal cell carcinoma or a non-invasive/in situ cervical cancer)
- 5. Has had certain other recent treatment e.g. major surgery, field radiation, received investigational agent, within the specified time frames prior to study drug administration 6. History of allergic reactions/hypersensitivity to bevacizumab (for patient who will receive
- bevacizumab) or any components used in the formulation
- 7. Grade 3 or higher hypersensitivity reaction to oxaliplatin, or grade 1-2 hypersensitivity reaction to oxaliplatin not controlled with premedication
- 8. Patient previously treated by S 95005 or history of allergic reactions attributed to compounds of similar or biologic composition to S 95005
- 9. Certain serious illnesses or serious medical conditions

10. Any condition that, in the judgment of the Investigator, may affect the patient's ability to understand and sign the informed consent and fully comply with all study procedure 11. Pregnancy or breast feeding

# Date of first enrolment 09/05/2016

Date of final enrolment 24/01/2019

# Locations

# Countries of recruitment United Kingdom England Austria

Germany

France

Hungary

Italy

Spain

Study participating centre Vall d'Hebron University Hospital (Hospital Vall D'Hebron) [VHIO] Passeig de la Vall d'Hebron, 119-129

Barcelona Spain 08035

Study participating centre University Hospital of Valencia (Hospital Clínic Universitari de València)

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Study participating centre

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# Study participating centre

Istituto Scientifico Romagnolo per lo Studio e la Cura dei Tumori (I.R.S.T.)

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# Study participating centre Klinikum der Universität München

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# Study participating centre

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# Study participating centre Universitätsklinikum Hamburg

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# Study participating centre Medizinische Universität Wien

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# Sponsor information

# Organisation

Servier (France)

### **ROR**

https://ror.org/034e7c066

# Funder(s)

# Funder type

Industry

# **Results and Publications**

# Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study will be available upon request from https://clinicaltrials.servier.com/ after the Marketing Authorisation has been granted.

# IPD sharing plan summary

Available on request

# **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Basic results			17/05/2021	No	No
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes
Plain English results			23/06/2021	No	Yes