Pancreatic cancer (PC) is an aggressive malignancy and the 4th cause of all cancer deaths worldwide. More than 30% of patients with PC are unresectable because of the local extension with a median overall survival of less than one year. The standard of care remains gemcitabine alone for unresectable locally advanced PC.

It was demonstrated that FOLFIRINOX (5-FU, irinotecan, oxaliplatin) is superior to gemcitabine in the treatment of resected PC, in terms of OS and progression-free survival (PFS). The recent PRODIGE 24 study showed that FOLFIRINOX is superior to gemcitabine in advanced therapy of resected PC, in terms of disease-free survival and OS. The superiority of FOLFIRINOX vs gemcitabine remains to be proven in unresectable locally advanced PC.

The aim of NEOPAN study is to evaluate the efficacy of FOLFIRINOX versus gemcitabine in the treatment of unresectable locally advanced PC.

Objectives

Primary objective: To compare progression-free survival (PFS) in patients with locally advanced adenocarcinoma of pancreas treated by Folfirinox or Gemcitabine

Secondary objectives:
- To compare the following endpoints between the 2 treatment arms:
  - Composite index of induction treatment severity toxicity (biliary tract infection grade 3-4 + any grade 3 toxicities + induction chemotherapy interruption rate) after 8 weeks of treatment.
  - Serious adverse events (NCI-CTC v4.0), compliance to treatment
  - Overall survival
  - Rate of events from PFS
  - Rate of surgery with curative intent (patients with R0 tumour resection)
  - Objective response rate and disease control rate (RECIST 1.1 criteria)
  - Time to treatment failure
  - Quality of life (ELO C30)

Study design

NEOPAN is a multicenter, phase 3, open-label randomised study (NCT02539537).

Patients treatment

Study treatment:
- Patients will be treated as follows during 6 months:
  - Control arm: Gemcitabine 1 000 mg/m² IV infusion over 30 min once per week, 3 weeks on/1 week off except for the first cycle with 4 infusions at D1, D8, D15, D22, repeated every 4 weeks for 6 cycles;
  - Experimental arm (mFolfirinox): leucovorin 400 mg/m² at D1, irinotecan 180 mg/m² at D1, oxaliplatin 85 mg/m² at D1, 5FU 2400 mg/m² at D3, oxaliplatin infusion over 4h, no bolus of 5FU, repeated every 2 weeks for 12 cycles.

Treatment recommendation after end of study treatment:
- After this 6-month period, each center has to choose between 3 strategies: stop of treatment, maintenance chemotherapy with gemcitabine or capcitabine, radio-chemotherapy.

Statistical considerations

Sample size: 170 patients, 142 events, 5 years of accrual and follow-up duration of at least 6 months

Based on assumptions of PFS prolongation from 5 months (control arm) to 8 months (experimental arm), a risk 3% (two-sided), power 80%.

Interim analyses:
- 4 interim analyses planned for the composite index of induction treatment severity toxicity to control Folfirinox tolerance after 8 weeks of treatment (4 cycles)

Study Population

Main Inclusion Criteria
- Patients ≥ 18 year of age
- Cytologically or histologically proven adenocarcinoma of the pancreas
- Locally advanced PC proven to be unresectable after multidisciplinary discussion
- Performance status ECOG < 2
- Normal hematologic, hepatic, and renal functions
- Adequate other vital functions
- Measurable lesions (RECIST criteria)
- Signed informed consent

Main Exclusion Criteria
- Metastatic disease
- Treatment of another cancer within 5 years prior to recruitment, except for in situ cervical cancer and basal cell carcinoma
- Cardiac disorders grade IV according to NYHA
- Major comorbidity (HIV, chronic HBV or HCV, non controlled diabetes)
- Pre-existing neuropathy grade ≥ 2, Gilbert disease or UCTA1 *28*P28
- Pregnancy

Recruiting sites

For additional information, please contact Prof. Michel DUCREUX, study coordinator: Michel.DUCREUX@gustaveroussy.fr

Acknowledgment

We thank the patients and their families for participating in the study. We are also indebted to all the participating centers.

Study Status

- Study timelines: Start of recruitment in Mar 2015
- Planned end of recruitment in March 2020
- Primary endpoint analysis expected in March 2021

IDMC meeting

- First IDMC in Jan 2018: Folfirinox toxicity was manageable and acceptable. Recommendation to continue the study without any change
- Next IDMC planned in Q1 2019

Accrual status

As of September 28th, 2018, 97 patients were identified over 27 centres. Among them 85 were randomized (Figure 3).

Figure 1 - NEOPAN study design

Figure 2 - Repartition of participating sites

Figure 3 - accrual status

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