

BMJ Open is committed to open peer review. As part of this commitment we make the peer review history of every article we publish publicly available.

When an article is published we post the peer reviewers' comments and the authors' responses online. We also post the versions of the paper that were used during peer review. These are the versions that the peer review comments apply to.

The versions of the paper that follow are the versions that were submitted during the peer review process. They are not the versions of record or the final published versions. They should not be cited or distributed as the published version of this manuscript.

BMJ Open is an open access journal and the full, final, typeset and author-corrected version of record of the manuscript is available on our site with no access controls, subscription charges or pay-per-view fees (<u>http://bmjopen.bmj.com</u>).

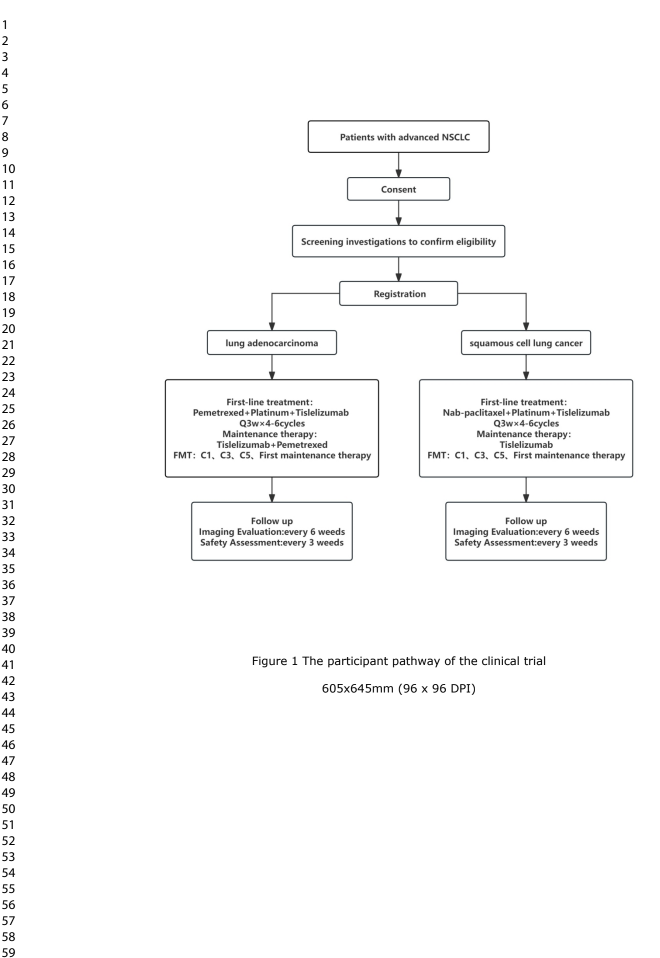
If you have any questions on BMJ Open's open peer review process please email <u>info.bmjopen@bmj.com</u>

# **BMJ Open**

# FMT combined with first-line treatment for driver-gene negative advanced NSCLC: study protocal for a prospective, multicenter, single-arm exploratory trial

Journal:	BMJ Open
Manuscript ID	bmjopen-2024-094366
Article Type:	Protocol
Date Submitted by the Author:	29-Sep-2024
Complete List of Authors:	Wei, Yanshuang; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology Qin, Lanqun; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Wu, Xinyu; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology Li, Dongqing; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Qian, Danping; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Geng, Qian; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Geng, Qian; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Jiang, Hua; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Jiang, Hua; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center
Keywords:	CHEMOTHERAPY, Clinical Relevance, Clinical Trial, ONCOLOGY, Respiratory tract tumours < ONCOLOGY





# FMT combined with first-line treatment for driver-gene negative advanced NSCLC: study protocal for a prospective, multicenter, single-arm exploratory trial

Yanshuang Wei<sup>1</sup><sup>†</sup>, Lanqun Qin<sup>1,2</sup><sup>†</sup>, Xinyu Wu<sup>1</sup>, Dongqing Li<sup>1,2</sup>, Danping Qian<sup>1,2</sup>, Qian Geng<sup>1,2\*</sup>, Hua Jiang<sup>1,2\*</sup>

# (†These authors contributed equally to this work and share first authorship)

<sup>1</sup>Department of Oncology, the Second Peoples' Hospital of Changzhou, the Third Affiliated Hospital of Nanjing Medical University, Changzhou, 213000, China. <sup>2</sup>Changzhou Medical Center, Nanjing Medical University, Changzhou, 213003, China.

# \*Correspondence:

# (Qian Geng is the main correspondence author and Hua Jiang is the secondary correspondence author)

Qian Geng, E-mail:karengq@njmu.edu.cn, Address: Department of Oncology, the Second Peoples' Hospital of Changzhou, the Third Affiliated Hospital of Nanjing Medical University, Changzhou, China.

Hua Jiang, E-mail:czeyjh@njmu.edu.cn, Address: Department of Oncology, the Second Peoples' Hospital of Changzhou, the Third Affiliated Hospital of Nanjing Medical University, Changzhou, China.

# ABSTRACT

**Introduction:** Chemotherapy combined with immunotherapy is the standard first-line treatment for driver-gene negative advanced non-small cell lung cancer (NSCLC). However, due to the immune microenvironment imbalance and immune status impairment caused by repeated chemotherapy, as well as the primary or secondary resistance to immune checkpoint inhibitors, the efficacy of immunotherapy combined with chemotherapy remains unsatisfactory. Recent studies have shown that fecal microbiota transplantation (FMT) can modulate the intestinal microflora, affecting the tumor immune microenvironment, and further enhancing the efficacy of tumor immunotherapy.

**Methods and analysis:** FMT-JSNO-02 (NCT06403111) is a prospective, multicenter, single-arm exploratory study. It is planned to include 62 cases of previously untreated driver-gene negative, ECOG PS 0-1, PD-L1<50% advanced NSCLC patients, who will

A word count: 3205

be given FMT by orally ingested stool capsules on the basis of first-line treatment of chemotherapy combined with immunotherapy. The safety and efficacy of this treatment regimen will be evaluated, with the primary endpoint being the 12-months progression-free survival (PFS) rate. Ethics and dissemination: Approved by the Ethics Committee of Changzhou No.2 People's Hospital (number [2024] YLJSA005). Trial registration number: NCT06403111. Strengths and limitations of this study: This is a multicenter prospective study. Our study aims to regulate intestinal flora and introduce FMT into the first-line treatment of lung cancer for the first time. The sample size in this study is relatively small. This is a single-arm study rather than a randomized controlled study. keywords: Fecal microbiota transplantation, Non Small Cell Lung Cancer, Chemotherapy, Immunotherapy 

For peer review only - http://bmjopen.bmj.com/site/about/guidelines.xhtml

BMJ Open: first published as 10.1136/bmjopen-2024-094366 on 4 March 2025. Downloaded from http://bmjopen.bmj.com/ on June 12, 2025 at Agence Bibliographique de Protected by copyright, including for uses related to text and Enseignement Superieur (ABES) ata mining, Al training, and similar technologies

BMJ Open: first published as 10.1136/bmjopen-2024-094366 on 4 March 2025. Downloaded from http://bmjopen.bmj.com/ on June 12, 2025 at Agence Bibliographique de Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

# 1. Introduction

# **Current Situation and Dilemmas of Immunotherapy for NSCLC**

Lung cancer is the most common malignancy worldwide and remains the leading cause of cancer-related mortality, accounting for 18.7% of all cancer deaths.<sup>[1]</sup>As the main pathological subtype of lung cancer, the majority of NSCLC patients present with local spread or distant metastasis upon diagnosis, losing the opportunity for surgery and having a very poor prognosis. In recent years, immune checkpoint inhibitors (ICIs), particularly those targeting the programmed death receptor 1 (PD-1) and its ligand (PD-L1), markedly improved the treatment outcomes and becoming an effective treatment strategy for NSCLC.

At present, the combination of chemotherapy and immunotherapy has become the standard first-line treatment for patients with advanced NSCLC due to extensive clinical studies conducted on Pembrolizumab and Nivolumab. Tislelizumab developed in China has also shown remarkable efficacy. In patients with lung adenocarcinoma, the addition of Tislelizumab to chemotherapy resulted in significantly prolonged median progression-free survival (mPFS) compared to chemotherapy alone (9.7 versus 7.6 month), with a 12-month PFS rate of 31.3%.<sup>[2]</sup> Similarly, in patients with squamous cell lung cancer, the mPFS for Tislelizumab plus chemotherapy group was also higher than that of chemotherapy alone (7.6 months versus 5.5 months).<sup>[3]</sup>

Although ICIs significantly improved the survival of patients with advanced NSCLC, primary or acquired immune resistance inevitably occurs in clinical practice. Existing research suggests that primary immune resistance is related to the dysfunction in tumor antigen processing and presentation, insufficient T cell infiltration in the tumor microenvironment (TME), and over expression of suppressive immune cells.<sup>[4]</sup> Acquired immune resistances is related to adaptive changes that occur in tumor cells and the TME

during exposure to immunotherapy.<sup>[5]</sup>

Nowadays, a growing body of research suggests that gut flora may be able to reverse immune resistance. For instance, gut microbiota can regulate the infiltration of immune cells in the TME by modulating innate and adaptive immunity. They can also reprogram the TME through their reactive metabolites or secretions, affecting the efficacy of ICIs and even reversing immune resistance.<sup>[6]</sup> Bifidobacterium can promote the maturation of dendritic cells and increase the infiltration of CD8<sup>+</sup> T cells in the TME as well.<sup>[7]</sup>

Similarily, research on the improvement of tumor immunotherapy efficacy by gut microbiota has been widely verified. Zitvoge found that the antitumor effects of cytotoxic T-lymphocyte-associated protein 4 (CTLA-4) blockade depended on distinct Bacteroides species. Tumors in antibiotic-treated or germ-free mice did not respond to CTLA-4 blockade. This defect was overcome by gavage with B. fragilis, by immunization with B. fragilis polysaccharides, or by adoptive transfer of B. fragilis-specific T cells.<sup>[8]</sup> At the same time, Sivan et al. found that mice orally administered Bifidobacterium could achieve the same anti-tumor effect as PD-L1 monoclonal antibodies, and the combination almost completely inhibited tumor growth.<sup>[7]</sup> Therefore, modulating gut microbes has the potential to further improve the treatment outcomes for patients receiving immunotherapy.

#### **Current Status of Fecal Microbiota Transplantation (FMT)**

The gut microbiota is susceptible to various factors, including the use of broad-spectrum antibiotics, repeated chemotherapy and other treatments. Common methods to regulate the intestinal microbiota include dietary intervention, probiotics, prebiotics, and FMT. Among these approaches, FMT stands as the most established technique.<sup>[9]</sup> FMT is the transplantation of functional flora from the stool of a healthy person into the intestine of a

Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies

BMJ Open: first published as 10.1136/bmjopen-2024-094366 on 4 March 2025. Downloaded from http://bmjopen.bmj.com/ on June 12, 2025 at Agence Bibliographique de

**Enseignement Superieur (ABES)** 

Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies

patient in a certain way. This method mainly includes two ways: injecting microflora fluids and swallowing capsules. Among them, FMT by orally ingested stool capsules (capsulized FMT) has the characteristics of a wide range of application, high acceptance degree, relative economy and convenience.<sup>[10]</sup>

FMT has been shown to ugment the antitumor effect of ICIs and overcome resistance to immunotherapy. In a study focused on 15 melanoma patients resistant to anti-PD-1 therapy, rechallenge with FMT and Pembrolizumab showed an objective responses in 3 out of 15 patients, and these 3 patients kept stable disease (SD) for more than 12 months.<sup>[11]</sup> In a clinical trial (NCT04264975), researchers performed FMT with anti-PD-1 inhibitors in 13 patients with anti-PD-1 refractory advanced solid cancers. The results showed that FMT induced sustained microbiota changes and clinical benefits in 6 patients, achieving an objective response rate (ORR) of 7.7% and a disease control rate (DCR) of 46.2%.<sup>[12]</sup> As a result, FMT holds great potential to reverse immune resistance and enhance the efficacy of immunotherapy.

Hence, we conduct such a prospective, exploratory study in order to verify whether FMT combined with first-line treatment can prolong the treatment efficacy for driver-gene negative NSCLC. By observing the adverse events and measuring the 12-months PFS rate, the safety and efficacy of this therapy will be verified. It may be possible to provide a more effective treatment option for NSCLC patients.

#### 2. Methods

# 2.1 Study Design

This is a prospective, multicenter, single-arm exploratory study aimed at evaluating the safety and efficacy of FMT combined with first-line treatment for driver-gene negative NSCLC patients. A total of 62 patients with driver-gene negative, ECOG PS 0-1, and

#### **BMJ** Open

PD-L1<50% who have not received prior treatment are expected to be included in this study without grouping. Patients with squamous cell lung cancer will receive Nab-paclitaxel + Platinum + Tislelizumab for 4-6 cycles and then Tislelizumab maintenance therapy. Patients with lung adenocarcinoma will receive Pemetrexed + Platinum + Tislelizumab for 4-6 cycles and then Tislelizumab + Pemetrexed maintenance therapy. All the enrolled patients or their designated agents will sign the informed consent form within 3 days before starting treatment. The study was approved by the ethics committee of Changzhou No.2 People's Hospital (number [2024] YLJSA005), and was registered at ClinicalTrials.gov (NCT06403111). See Figures 1 for details

# 2.2 Patient Population

# **Inclusion criteria:**

- The subjects voluntarily joined the study and were able to sign the informed consent with good compliance.
- Age 18-80 years old (when signing the informed consent form).
- Patients with histologically or cytologically proven locally advanced (III B/III C), metastatic, or recurrent (stage IV) NSCLC who are inoperable and unable to receive radical concurrent chemoratherapy, according to the International Association for the Study of Lung Cancer and the American Joint Committee on Cancer Classification, 8th Edition TNM Classification of Lung cancer.
- Have not received systemic intravenous anti-tumor therapy before, and the driver-gene is negative.
- PD-L1 expression <50%.
- According to the solid tumor efficacy evaluation criteria (RECIST 1.1), there is at least one radiographically measurable lesion. That is, in CT or MRI detection, the longest diameter of a single lesion was ≥10mm, or the pathological enlargement of a single lymph node was ≥15mm.
- The physical status score of Eastern Tumor Collaboration Group (ECOG) was 0-1.

BMJ Open: first published as 10.1136/bmjopen-2024-094366 on 4 March 2025. Downloaded from http://bmjopen.bmj.com/ on June 12, 2025 at Agence Bibliographique de

Enseignement Superieur (ABES)

Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies

• Expected survival >3 months.

- Have adequate organ and bone marrow function, laboratory tests within 7 days prior to enrollment meet the following requirements (no blood components, cell growth factors, albumin and other corrective drugs are allowed within 14 days prior to obtaining laboratory tests), as follows: 1) Blood routine: absolute neutrophil count (ANC)  $\geq 1.5 \times 109/L$ , platelet (PLT)  $\geq 75 \times 109/L$ , hemoglobin (HGB)  $\geq 90$  g/L (no blood transfusion or erythropoietin dependence within 14 days). 2) Liver function: serum total bilirubin (TBIL)  $\leq 2$  times the upper limit of normal (ULN). Alanine aminotransferase (ALT) and/or aspartate aminotransferase (AST)  $\leq$  5x ULN, serum albumin  $\geq 28$  g/L. alkaline phosphatase (ALP)  $\leq 5 \times ULN$ . 3) Renal function: serum creatinine (Cr)  $\leq 1.5 \times ULN$ , or creatinine clearance  $\geq 50$  mL/min (using the standard Cockcroft-Gault formula) : Urine routine results showed urinary protein < 2+. For patients with urine protein  $\geq 2^+$  at baseline, 24-hour urine collection and 24-hour urine protein quantification < 1g should be performed. 4) Coagulation function: International standardized ratio (INR) or prothrombin time (PT)  $\leq 1.5$  times ULN. If the subject is receiving anticoagulant therapy, as long as the INR is within the intended range of anticoagulant drug use.
- For female subjects of reproductive age, a urine or serum pregnancy test should be performed and the result is negative 3 days prior to receiving the initial study drug administration.
- Subjects and their sexual partners are required to use a medically approved contraceptive method (such as an intrauterine device (IUD), contraceptive pill, or condom) during the study treatment period and for 6 months after the end of the study treatment period.

### **Exclusion criteria:**

• Currently participating in an interventional clinical study or receiving another investigational drug or investigational device within 4 weeks prior to initial dosing.

#### **BMJ** Open

- Received proprietary Chinese medicines with anti-tumor indications or immunomodulatory drugs (thymosin, interferon, interleukin, etc.) within 2 weeks before the first administration, or received major surgical treatment within 3 weeks before the first administration.
  - Class III IV congestive heart failure (New York Heart Association classification), poorly controlled and clinically significant arrhythmias.
  - Any arterial thrombosis, embolism or ischemia, such as myocardial infarction, unstable angina pectoris, cerebrovascular accident or transient ischemic attack, occurred within 6 months before treatment.
  - Known allergic reaction to the drug in this study.
  - Patients requiring long-term systemic use of corticosteroids. Patients with chronic obstructive pulmonary disease (COPD) or asthma requiring intermittent use of bronchodilators, inhaled corticosteroids, or local corticosteroids could be enrolled.
  - Symptomatic central nervous metastases. Patients with asymptomatic brain metastases (BMS) or BMS whose symptoms are stable after treatment are eligible to participate in this study if they meet all of the following criteria: measurable lesions outside the central nervous system. No midbrain, pontine, cerebellum, meninges, medulla oblongata or spinal cord metastasis. Maintain clinical stability for at least 2 weeks. Stop hormone therapy 3 days before the first dose of the study drug.
  - There is an active infection requiring treatment or systemic anti-infective drugs have been used in the week prior to the first dosing.
  - Has not fully recovered from toxicity and/or complications caused by any intervention before starting treatment (i.e., ≤ grade 1 or baseline, excluding weakness or hair loss).
  - Known history of human immunodeficiency virus (HIV) infection (i.e. HIV 1/2 antibody positive).
  - Untreated active hepatitis B (defined as HBsAg positive and HBV-DNA copy

number detected greater than the upper limit of normal value in the laboratory of the study center). Note: Hepatitis B subjects who meet the following criteria can also be enrolled: HBV viral load <1000 copies /ml (200 IU/ml) before initial administration, subjects should receive anti-HBV therapy throughout the study chemotherapy drug treatment to avoid viral reactivation. 2) For subjects with anti-HBC (+), HBsAg (-), anti-HBS (-) and HBV viral load (-), prophylactic anti-HBV therapy is not required, but close monitoring of viral reactivation is required.

- Active HCV-infected subjects (HCV antibody positive and HCV-RNA levels above the lower limit of detection).
- Received live vaccine within 30 days prior to the first dose (cycle 1, day 1). Note: Injectable inactivated virus vaccine against seasonal influenza is permitted for 30 days prior to initial administration. However, live attenuated influenza vaccines administered intranasally are not permitted.
- Pregnant or lactating women.

• Medical history or evidence of disease that may interfere with test results, prevent participants from fully participating in the study, abnormal treatment or laboratory test values, or other conditions that the investigator considers unsuitable for enrollment. The Investigator considers other potential risks unsuitable for participation in the study.

# 2.3 Treatment Regimen

Dose selection: Tislelizumab: 200mg. Pemetrexed: 500mg/m<sup>2</sup>. Nab-paclitaxel: 260mg/m<sup>2</sup>. Platinum drugs (Cisplatin: 60-80mg/m<sup>2</sup>; Carboplatin: AUC=5; Nedaplatin: 80mg/m<sup>2</sup>). Capsulized FMT: 30 capsules each time. Capsules will be stored in the refrigerator at -80° C. When taking capsules, take them out of the refrigerator, heat them in a water bath (single-hole water bath, JF-01 defroster) at 37 °C for 10 minutes, and swallow them with warm water.

Patients will receive the first-line regimen of Nab-paclitaxel + Platinum + Tislelizumab

(squamous cell lung cancer) / Pemetrexed + Platinum + Tislelizumab (lung adenocarcinoma) for 4-6 cycles. If there is no progression of the disease after 4-6 cycles of first-line treatment, then patients will enter the maintenance treatment stage. Patients will receive Tislelizumab maintenance treatment (squamous cell lung cancer ), or Tislelizumab + Pemetrexed maintenance treatment (lung adenocarcinoma). Capsulized FMT should be conducted on the 5th day ( $\pm 2$  days) after the start of the 1, 3 and 5 cycles of chemotherapy, as well as on the 5th day ( $\pm 2$  days) after the first maintenance chemotherapy (lung adenocarcinoma) and before the first maintenance immunotherapy (squamous cell lung cancer). See Figures 2 for details

# 2.4 Study Endpoints

The primary endpoint of the study is the 12-month PFS rate. Secondary endpoints include ORR, mPFS, overall survival (OS), duration of response (DOR), safety, microbiome diversity, and quality of life (QoL). Exploratory endpoints including efficacy predictive biomarkers, including fungi, bacteria, metabolomics, and proteomics.

# 2.5 Sample Size Estimation

The parameters are set as follows: the two-sided significance level alpha is set at 0.05. Based on an estimated dropout rate of 0% and a reference 12-month PFS rate of 35% published in the Rational-304 and 307 clinical studies, it is inferred that the 12-month PFS rate in this trial can reach 55%.<sup>[2,3]</sup> Therefore, a total sample size of 62 participants is determined for this study to ensure statistical power greater than 80%.

# 2.6 Efficacy and Safety Assessments Imaging evaluation

The assessments of the screening period should be conducted within 28 days prior to the first administration of the study drug. Prior to treatment, researchers at the study center will confirm that subjects have measurable lesions that met the RECIST 1.1 criteria. The methods used to assess tumor burden at baseline must be consistent with those used for each subsequent follow-up assessment (CT/MRI). Additional imaging assessments of

other suspected involved areas (eg, brain) may be conducted based on the subject's clinical symptoms and signs. After enrollment, tumor status will be evaluated using imaging methods every 6 weeks ( $\pm$ 7 days), every 12 weeks ( $\pm$ 7 days) after 48 weeks until disease progression (RECIST 1.1) or death (during the course of patient treatment). For subjects who have completed treatment or have discontinued treatment for reasons other than disease progression, a single tumor imaging assessment should be performed at the time of treatment completion/discontinuation.

# Safety evaluation

Researchers will conduct safety assessments on patients every 3 weeks using NCI-CTCAE 5.0. After treatment ends, participants will be followed up for 30 days to detect adverse events (AE). If the patient has not received new anti-tumor treatment within 90 days after the last dose, severe adverse events (SAE) occurring within 90 days after the last dose, severe adverse events (SAE) occurring within 90 days after the last dose will be collected. If the subject has received new anti-tumor treatment, SAE prior to the new treatment will be collected, with precedence given to those that have already occurred. Investigators should grade and record adverse events for each subject according to the NCI-CTCAE 5.0 criteria during the study and follow-up period. The characteristics of adverse events will be assessed and recorded based on severity, causality, toxicity grading, management measures, and outcomes.

#### Fecal and peripheral blood collection time

With the permission of the ethics committee, patients should provide 10ml whole blood samples and fecal samples at baseline, after two cycles of treatment, before maintenance treatment, and after two cycles of maintenance treatment for the detection of efficacy prediction markers (each cycle is 21 days).

# 2.7 Data Management Data management methods

The study will use a data collection system (91trial) for data management. All subjects will be assigned a unique ID. Researchers will input basic information, drug use, laboratory examination and other raw data information of subjects into 91trial. The system is subject to superior monitoring and cannot be arbitrarily modified. The data manager will write a data audit report based on the trial protocol and audit criteria in the database. The results of the patient's report on tests, examinations, etc. can be obtained from the electronic case report. Quality of life will be assessed using the EORTC-QLQ-C30 questionnaire, which will be available in both paper and electronic formats.

# Reporting and collection of adverse events

When a clinical adverse event occurs, it will be detailed on the case report form with the time of occurrence, clinical manifestations, course of treatment, duration, outcome, and the relationship to this study. For those with laboratory abnormalities, follow-up will continue until the test results return to normal, or to the level before medical intervention, or until it is determined that the event is unrelated to the medical intervention. In the event of a serious adverse event, the serious adverse event form should be filled out and reported to the department and relevant functional departments within 24 hours and recorded in the hospital quality control system (HQS) system, with documentation in the medical record.

### 2.8 Statistical Analysis

SPSS Statistic will be used for statistical processing in this study. Kaplan-Meier survival analysis will be used to draw PFS and OS survival curves, and Log-rank or Breslow test will be used to explore statistical significance. Multivariate analysis of PFS rate and OS will be performed by Cox proportional risk model.

### 2.9 Patient and public involvement

Patients and/or the public were not involved in the design, or conduct, or reporting, or

BMJ Open: first published as 10.1136/bmjopen-2024-094366 on 4 March 2025. Downloaded from http://bmjopen.bmj.com/ on June 12, 2025 at Agence Bibliographique de Enseignement Superieur (ABES)

Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies

dissemination plans of this research.

# 3. Ethics and Dissemination

This prospective study has been approved by the Clinical Medical Technology Ethics Committee of Changzhou No.2 People's Hospital (number [2024] YLJSA005) and has been registered in ClinicalTrials.gov (NCT06403111).

# 4. Discussion

The application of ICIs has brought significant benefits to patients with driver gene-negative NSCLC, but statistics show that the 5-year survival rate for advanced NSCLC patients receiving immunotherapy is only about 20%.<sup>[13]</sup> With a deeper exploration of the mechanisms of immune resistance, researchers have found that the gut microbiota plays a crucial role in regulating the immune system and can significantly influence the effectiveness of immunotherapy.

Preclinical trials have demonstrated that specific gut microbiota or fecal microbiota transplantation using feces from patients who respond to ICIs can modulate the immune system, enhance immune cell infiltration in tumors, induce tumor regression, and improve the antitumor efficacy of ICIs.<sup>[7,14–16]</sup> Existing clinical trials have shown that exogenous microflora supplementation by FMT can change the composition of intestinal microflora and the proportion of dominant microflora, reprogram the tumor microenvironment, and overcome immune resistance.<sup>[11,17]</sup> However, the majority of related studies have focused on melanoma, with limited research in the field of lung cancer treatment. A proof-of-concept clinical trial conducted by a team of researchers in Korea has demonstrated the potential benefits of FMT in a clinical setting other than melanoma, giving us confidence to carry out our study.<sup>[12]</sup> Therefore, we designed a multi-prospective, multicenter, single-arm exploratory study of patients with advanced NSCLC. If patients treated with FMT have longer PFS and higher 12-months PFS rate,

this study is expected to provide strong evidence for further improving outcomes in lung cancer patients.

# **Clinical Trial Status**

At the time of submission, the study is ongoing and open to recruitment patients.

# **Authors' Contributions**

YW and LQ are responsible for the article's concept and writing; QG, DL, and DQ are responsible for clinical treatment; QG, LQ, YW, and XW are responsible for data collection and compilation. HJ and QG coordinate all the work. All authors review and approve the final manuscript.

# Funding

This study was supported by the 2022 Clinical Research project of Changzhou Medical Center, Nanjing Medical University (CMCC202201), 2022 Changzhou 8th Batch of Science and Technology Project (Applied Basic Research) (CJ20220086).

# **Competing interests**

All authors have completed the ICMJE uniform disclosure form at http://www.icmje.org/coi\_disclosure.pdf and declare: no support from any organization for the submitted work; no financial relationships with any organization that might have an interest in the submitted work in the previous three years, no other relationships or activities that could appear to have influenced the submitted work.

# Acknowledgements

We would like to express our sincere appreciation to all the participants.

# Peer review

Not externally peer reviewed.

# Data sharing

No additional data are available.

# References

1 Beay F, Laversanne M, Sung H, et al. Global cancer statistics 2022: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. *CA Cancer J Clin* 2024;74(3):229-263 doi:10.3322/caac.21834 [published Online First: 4 April 2024].

2 Lu S, Wang J, Yu Y, et al. Tislelizumab Plus Chemotherapy as First-Line Treatment for Locally Advanced or Metastatic Nonsquamous NSCLC (RATIONALE 304): A Randomized Phase 3 Trial. *J Thorac Oncol* 2021;16(9):1512-1522 doi:10.1016/j.jtho.2021.05.005 [published Online First: 23 May 2021].

3 Wang J, Lu S, Yu X, et al. Tislelizumab Plus Chemotherapy vs Chemotherapy Alone as First-line Treatment for Advanced Squamous Non–Small-Cell Lung Cancer: A Phase 3 Randomized Clinical Trial. *JAMA Oncol* 2021;7(5):709-717 doi:10.1001/jamaoncol.2021.0366.

4 Nagasaki J, Ishino T, Togashi Y. Mechanisms of resistance to immune checkpoint inhibitors. *Cancer Sci* 2022;113(10):3303-3312 doi:10.1111/cas.15497 [published Online First: 30 July 2022].

5 Bagchi S, Yuan R, Engleman EG. Immune Checkpoint Inhibitors for the Treatment of Cancer: Clinical Impact and Mechanisms of Response and Resistance. *Annu Rev Pathol* 2021;16:223-249 doi:10.1146/annurev-pathol-042020-042741 [published Online First: 16 November 2020].

6 Wu J, Wang S, Zheng B, Qiu X, et al. Modulation of Gut Microbiota to Enhance Effect of Checkpoint Inhibitor Immunotherapy. *Front Immunol* 2021;12:669150 doi:10.3389/fimmu.2021.669150 [published Online First: 29 June 2021].

7 Sivan A, Corrales L, Hubert N, et al. Commensal Bifidobacterium promotes antitumor immunity and facilitates anti-PD-L1 efficacy. *Science* 2015;350(6264):1084-1089.DOI:10.1126/science.aac4255 [published Online First: 5 November 2015].

8 Vétizou M, Pitt JM, Daillère R, et al. Anticancer immunotherapy by CTLA-4 blockade relies on the gut microbiota. *Science* 2015;350(6264):1079-1084 doi:10.1126/science.aad1329 [published Online First: 5 November 2015].

9 Blake SJ, Wolf Y, Boursi B, et al. Role of the microbiota in response to and recovery from cancer therapy. *Nat Rev Immunol* 2024;24(5):308-325 doi:10.1038/s41577-023-00951-0 [published Online First: 6 November 2023].

10 Erdmann J. How gut bacteria could boost cancer treatments. *Nature* 2022;607(7919):436-439 doi:10.1038/d41586-022-01959-7 [published Online First: 19 July 2022].

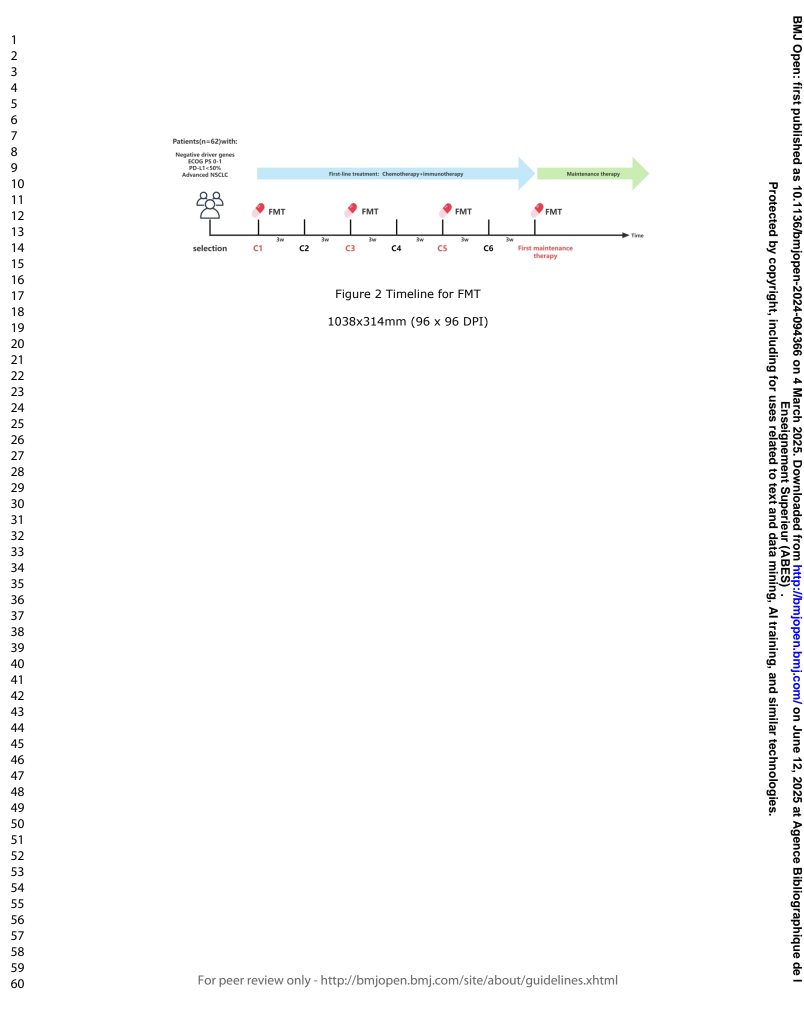
11 Davar D, Dzutsev AK, Mcculloch JA, et al. Fecal microbiota transplant overcomes resistance to anti–PD-1 therapy in melanoma patients. *Science* 2021; 371(6529): 595-602 doi:10.1126/science.abf3363 [published Online First: 5 February 2021].

12 Kim Y, Kim G, Kim S, et al. Fecal microbiota transplantation improves anti-PD-1 inhibitor efficacy in unresectable or metastatic solid cancers refractory to anti-PD-1 inhibitor. *Cell Host Microbe* 2024;32(8):1380-1393.e9 doi:10.1016/j.chom.2024.06.010 [published Online First: 25 July 2024].

13 Xing Li, Fei Zhou, Chunxia Su. Advances in the research of mechanisms and strategies in immunotherapy resistance. *Chinese Journal of Metastatic Cancer (in Chinese)* 2020,03(03):235-240 doi:10.3760/cma.j.cn101548-20200520-00060 [published Online First: 30 September 2020].

#### **BMJ** Open

• •	cience.abc3421 Le Chatelier E		t al. Gut mic		nfluences ef	•
immunothera doi:10.1126/s	py against cience.aan3706	epithelial [published On]	tumors. line First: 2 N	<i>Scienc</i> Jovember 2		359(6371
	shnan V, Spence				-	response to an
	by in melanoma	-	ice 2018; 359	9(6371): 97	-103 doi:10.	1126/science.a
	lline First: 2 Nov N, Youngster I, 1	-	G, et al. Feca	l microbio	ta transplant	promotes rest
immunothera	<u> </u>			a ·	2021 2	71(6529):
doi:10.1126/s	cience.abb5920	[published On	line First: 10	December	2020].	
		[published On				



# Fecal microbiota transplantation combined with platinumbased doublet chemotherapy and Tislelizumab as first-line treatment for driver-gene negative advanced non-small-cell lung cancer (NSCLC): study protocol for a prospective, multi-center, single-arm exploratory trial

Journal:	BMJ Open
Manuscript ID	bmjopen-2024-094366.R1
Article Type:	Protocol
Date Submitted by the Author:	20-Dec-2024
Complete List of Authors:	<ul> <li>Wei, Yanshuang; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology</li> <li>Qin, Lanqun; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center</li> <li>Wu, Xinyu; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology</li> <li>Li, Dongqing; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology</li> <li>Li, Dongqing; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center</li> <li>Qian, Danping; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center</li> <li>Jiang, Hua; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center</li> <li>Jiang, Hua; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center</li> <li>Geng, Qian; The Affiliated Changzhou No 2 People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center</li> </ul>
<b>Primary Subject Heading</b> :	Oncology
Secondary Subject Heading:	Research methods
Keywords:	CHEMOTHERAPY, Clinical Relevance, Clinical Trial, ONCOLOGY, Respiratory tract tumours < ONCOLOGY



2		
3	1	Fecal microbiota transplantation combined with platinum-based doublet
4	1	recar microbiota transplantation combined with platinum-based doublet
5 6	2	chemotherapy and Tislelizumab as first-line treatment for driver-gene negative
7 8	3	advanced non-small-cell lung cancer (NSCLC): study protocol for a prospective,
9 10	4	multi-center, single-arm exploratory trial
11		
12	5	Yanshuang Wei <sup>1</sup> <sup>+</sup> , Lanqun Qin <sup>1,2</sup> <sup>+</sup> , Xinyu Wu <sup>1</sup> , Dongqing Li <sup>1,2</sup> , Danping Qian <sup>1,2</sup> ,
13 14	6	Hua Jiang <sup>1,2</sup> *, Qian Geng <sup>1,2</sup> *
15		
16	7	(†These authors contributed equally to this work and share first authorship)
17		
18	8	<sup>1</sup> Department of Oncology, the Second Peoples' Hospital of Changzhou, the Third
19 20	9	Affiliated Hospital of Nanjing Medical University, Changzhou, 213000, China.
20	10	<sup>2</sup> Changzhou Medical Center, Nanjing Medical University, Changzhou, 213003, China.
22		
23	11	*Correspondence:
24	12	(Qian Geng is the main correspondence author and Hua Jiang is the secondary
25	13	correspondence author)
26 27		
28	14	Qian Geng, E-mail: karengq@njmu.edu.cn, Address: Department of Oncology, the Second Peoples'
29	15	Hospital of Changzhou, the Third Affiliated Hospital of Nanjing Medical University, Changzhou,
30	16	China.
31	10	
32	17	Hua Jiang, E-mail: czeyjh@njmu.edu.cn, Address: Department of Oncology, the Second Peoples'
33 34	18	Hospital of Changzhou, the Third Affiliated Hospital of Nanjing Medical University, Changzhou,
35		
36	19	China.
37	20	ABSTRACT
38	20	ADSTRACT
39 40	21	Introduction: Chemotherapy combined with immunotherapy is the standard first-line
41	22	treatment for driver-gene negative advanced non-small cell lung cancer (NSCLC).
42	23	However, due to the immune microenvironment imbalance and immune status
43		
44 45	24	impairment caused by repeated chemotherapy, as well as the primary or secondary
45 46	25	resistance to immune checkpoint inhibitors (ICIs), the efficacy of immunotherapy
47	26	combined with chemotherapy remains unsatisfactory. Recent studies have shown that
48	27	fecal microbiota transplantation (FMT) can modulate the intestinal microflora,
49	28	influence the tumor immune microenvironment, and even enhance the efficacy of
50 51	29	immunotherapy. Therefore, we conduct this study to evaluate the efficacy and safety of
51 52	30	FMT combined with standard first-line treatment for driver gene-negative advanced
53	31	NSCLC.
54	32	Methods and analysis: FMT-JSNO-02 (NCT06403111) is a prospective, multi-center,
55	33	single-arm exploratory study. It is planned to include 62 cases of previously untreated
56		
57 58	34	driver-gene negative, Eastern Cooperative Oncology Group Performance Status
50	35	(ECOG PS) 0-1, Programmed death ligand 1 (PD-L1) <50% advanced NSCLC

patients, who will be given FMT by orally ingested stool capsules on the basis of 

standard first-line treatment of chemotherapy combined with immunotherapy. The
safety and efficacy of this treatment regimen will be evaluated, with the primary

39 endpoint being the 12-months progression-free survival (PFS) rate.

Ethics and dissemination: The study was approved by the ethics committee of the
Second Peoples' Hospital of Changzhou (number [2024] YLJSA005) and is being
conducted in accordance with the principles of the Declaration of Helsinki. The results
of the study will be submitted for publication in peer-reviewed journals.

44 Trial registration number: ClinicalTrials.gov Identifier: NCT06403111. Date of
 45 registration: May 7, 2024, the first version protocol.

# 46 Strengths and limitations of this study:

- This is a multi-center prospective study.
- Our study aims to regulate intestinal flora and introduce FMT into the standard first-line treatment of lung cancer for the first time.
- The sample size in this study is relatively small.

• This is a single-arm study rather than a randomized controlled study.

keywords: Fecal microbiota transplantation, Non-Small Cell Lung Cancer,
 Chemotherapy, Immunotherapy

# **1. Introduction**

# 56 Current Situation and Dilemmas of Immunotherapy for Non-Small Cell lung 57 cancer (NSCLC)

Lung cancer is the most common malignancy worldwide and remains the leading cause of cancer-related mortality, accounting for 18.7% of all cancer deaths.<sup>[1]</sup>As the main pathological subtype of lung cancer, the majority of NSCLC patients present with local spread or distant metastasis upon diagnosis, losing the opportunity for surgery and having a very poor prognosis. In recent years, immune checkpoint inhibitors (ICIs), particularly those targeting the programmed death receptor 1 (PD-1) and its ligand (PD-L1), markedly improved the treatment outcomes and becoming an effective treatment strategy for NSCLC. 

At present, the combination of chemotherapy and immunotherapy has become the standard first-line treatment for patients with advanced NSCLC due to extensive clinical studies conducted on pembrolizumab and nivolumab. Tislelizumab developed in China has also shown remarkable efficacy. In patients with lung adenocarcinoma, Page 3 of 17

#### **BMJ** Open

the addition of tislelizumab to chemotherapy resulted in significantly prolonged median progression-free survival (mPFS) compared to chemotherapy alone (9.7 versus 7.6 month), with a 12-month PFS rate of 31.3%.<sup>[2]</sup> Similarly, in patients with lung squamous cell carcinoma, the mPFS for tislelizumab plus chemotherapy group was also higher than that of chemotherapy alone (9.6 months versus 5.5 months).<sup>[3]</sup>

Although ICIs significantly improved the survival of patients with advanced NSCLC, primary or acquired immune resistance inevitably occurs in clinical practice. Existing research suggests that primary immune resistance is related to the dysfunction in tumor antigen processing and presentation, insufficient T cell infiltration in the tumor microenvironment (TME), and over expression of suppressive immune cells.<sup>[4]</sup> Acquired immune resistances is related to adaptive changes that occur in tumor cells and the TME during exposure to immunotherapy.<sup>[5]</sup>

Nowadays, a growing body of research suggests that gut flora may be able to reverse immune resistance. For instance, gut microbiota can regulate the infiltration of immune cells in the TME by modulating innate and adaptive immunity. They can also reprogram the TME through their reactive metabolites or secretions, affecting the efficacy of ICIs and even reversing immune resistance.<sup>[6]</sup> Researchers have conducted preliminary explorations into how gut microbiota affects the efficacy of immunotherapy. Zitvoge found that the antitumor effects of cytotoxic T-lymphocyte-associated protein 4 (CTLA-4) blockade depended on distinct Bacteroides species. Tumors in antibiotic-treated or germ-free mice did not respond to CTLA-4 blockade. This defect was overcome by gavage with B. fragilis, by immunization with B. fragilis polysaccharides, or by adoptive transfer of B. fragilis-specific T cells.<sup>[7]</sup> At the same time, Sivan et al. found that mice orally administered Bifidobacterium could achieve the same anti-tumor effect as PD-L1 monoclonal antibodies, and the combination almost completely inhibited tumor growth.<sup>[8]</sup> Therefore, modulating gut microbes has the potential to further improve the treatment outcomes for patients receiving immunotherapy.

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

# 101 Current Status of Fecal Microbiota Transplantation (FMT)

The gut microbiota is susceptible to various factors, including the use of broad-spectrum antibiotics, repeated chemotherapy and other treatments. Common methods to regulate the intestinal microbiota include dietary intervention, probiotics, prebiotics, and FMT. Among these approaches, FMT stands as the most established technique.<sup>[9]</sup> FMT is the transplantation of functional flora from the stool of a healthy person into the intestine of a patient in a certain way. This method mainly includes two ways: injecting microflora fluids and swallowing capsules. Among them, FMT by orally ingested stool capsules (capsulized FMT) has the characteristics of a wide range of application, high acceptance degree, relative economy and convenience.<sup>[10]</sup> 

The relationship between intestinal flora and immunotherapy efficacy has been preliminarily confirmed in some clinical studies of malignant tumors. For example, in a study focused on 15 melanoma patients resistant to anti-PD-1 therapy, rechallenge with FMT and Pembrolizumab showed objective responses in 3 out of 15 patients, and these 3 patients kept stable disease (SD) for more than 12 months.<sup>[11]</sup> In a clinical trial (NCT04264975), researchers performed FMT with anti-PD-1 inhibitors in 13 patients with anti-PD-1 refractory advanced solid cancers. The results showed that FMT induced sustained microbiota changes and clinical benefits in 6 patients, achieving an objective response rate (ORR) of 7.7% and a disease control rate (DCR) of 46.2%.<sup>[12]</sup> As a result, FMT holds great potential to reverse immune resistance and enhance the efficacy of immunotherapy.

Hence, we conduct such a prospective, exploratory study in order to evaluate the efficacy and safety of FMT combined with standard first-line treatment for driver genenegative advanced NSCLC. We expect that the addition of FMT to the standard firstline treatment in NSCLC will be safe and will have an enhancing effect.

- 128 2. Methods
- **2.1 Study Design**

Page 5 of 17

#### **BMJ** Open

This is a prospective, multi-center, single-arm exploratory study aimed at evaluating the efficacy and safety of FMT combined with standard first-line treatment for driver-gene negative NSCLC patients. The trial protocol was written in accordance with the Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT) guidance. The study will mainly be conducted at the Second Peoples' Hospital of Changzhou, the Third Affiliated Hospital of Nanjing Medical University. Recruitment information can be obtained through outpatient consultations, posters displayed in the wards, and other channels. A total of 62 patients with driver-gene negative, Eastern Cooperative Oncology Group (ECOG) performance status (PS) 0-1, and PD-L1<50% who have not received prior treatment are expected to be included in this study without grouping. Patients will receive platinum-based doublet chemotherapy combined with tislelizumab as standard first-line treatment for 4-6 cycles and then enter maintenance therapy. FMT will be given to patients during the first, third, and fifth treatment cycles, as well as during the first maintenance therapy. All the enrolled patients or their designated agents will sign the informed consent form within 3 days before starting treatment. This study was approved by the ethics committee of the Second Peoples' Hospital of Changzhou (number [2024] YLJSA005). See figure 1 for details. 

- 147 2.2 Patient Population148 Inclusion criteria:
- The subjects voluntarily joined the study and were able to sign the informed
   consent with good compliance.

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

- Age 18-80 years old (when signing the informed consent form).
- Patients with histologically or cytologically proven locally advanced (IIIB/IIIC), metastatic, or recurrent (stage IV) NSCLC who are inoperable and unable to receive radical concurrent chemoradiotherapy, according to the International Association for the Study of Lung Cancer and the American Joint Committee on Cancer Classification, 8th Edition TNM Classification of Lung cancer.
- Have not received systemic intravenous anti-tumor therapy before, and the drivergene is negative.
- 159 PD-L1 expression < 50%.

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

BMJ Open

According to the solid tumor efficacy evaluation criteria (RECIST version 1.1),
 there is at least one radiographically measurable lesion. That is, in CT or MRI
 detection, the longest diameter of a single lesion was ≥ 10mm, or the pathological
 enlargement of a single lymph node was ≥ 15mm.

The physical status score of Eastern Tumor Collaboration Group (ECOG) was 0 1.

166 •

Expected survival > 3 months.

Have adequate organ and bone marrow function, laboratory tests within 7 days prior to enrollment meet the following requirements (no blood components, cell growth factors, albumin or other corrective drugs are allowed within 14 days prior to obtaining laboratory examination), as follows: 1) Blood routine: absolute neutrophil count (ANC)  $\geq 1.5 \times 10^{9}$ /L, platelet (PLT)  $\geq 75 \times 10^{9}$ /L, hemoglobin  $(HGB) \ge 90$  g/L (no blood transfusion or erythropoietin dependence within 14 days). 2) Liver function: serum total bilirubin (TBIL)  $\leq 2$  times the upper limit of normal (ULN). Alanine aminotransferase (ALT) and/or aspartate aminotransferase  $(AST) \le 5 \times ULN$ , serum albumin  $\ge 28$  g/L. alkaline phosphatase  $(ALP) \le 5 \times ULN$ . 3) Renal function: serum creatinine (Cr)  $\leq 1.5 \times ULN$ , or creatinine clearance  $\geq 50$ mL/min (using the standard Cockcroft-Gault formula): Urine routine results showed urinary protein < 2+. For patients with urine protein  $\ge 2+$  at baseline, 24-hour urine collection and 24-hour urine protein quantification < 1g should be performed. 4) Coagulation function: International standardized ratio (INR) or prothrombin time (PT)  $\leq$  1.5 times ULN. If the subject is receiving anticoagulant therapy, as long as the INR is within the intended range of anticoagulant drug use. For female subjects of reproductive age, a urine or serum pregnancy test should be performed and the result is negative 3 days prior to receiving the initial study drug administration. 

Subjects and their sexual partners are required to use a medically approved contraceptive method (such as an intrauterine device (IUD), contraceptive pill, or condom) during the study treatment period and for 6 months after the end of the study treatment period.

1 2		
3 4	190	Exclusion criteria:
5 6	191	• Currently participating in an interventional clinical study or receiving another
7 8	192	investigational drug or investigational device within 4 weeks prior to initial dosing.
9	193	• Received proprietary Chinese medicines with anti-tumor indications or
10 11	194	immunomodulatory drugs (thymosin, interferon, interleukin, etc.) within 2 weeks
12 13	195	before the first administration, or received major surgical treatment within 3 weeks
14 15	196	before the first administration.
16 17	197	• Class III - IV congestive heart failure (New York Heart Association classification),
18 19	198	poorly controlled and clinically significant arrhythmias.
20 21	199	• Any arterial thrombosis, embolism or ischemia, such as myocardial infarction,
22 23	200	unstable angina pectoris, cerebrovascular accident or transient ischemic attack,
24 25	201	occurred within 6 months before treatment.
26 27	202	• Known allergic reaction to the drug in this study.
28 29	203	• Patients who require long-term oral, intravenous, or intramuscular administration
30	204	of systemic corticosteroids.
31 32	205	• Symptomatic central nervous metastases. Patients with asymptomatic brain
33 34	206	metastases (BMS) or BMS whose symptoms are stable after treatment are eligible
35 36	207	to participate in this study if they meet all of the following criteria: measurable
37 38	208	lesions outside the central nervous system. No midbrain, pontine, cerebellum,
39 40	209	meninges, medulla oblongata or spinal cord metastasis. Maintain clinical stability
41 42	210	for at least 2 weeks. Stop hormone therapy 3 days before the first dose of the study
43 44	211	drug.
45 46	212	• There is an active infection requiring treatment or systemic anti-infective drugs
47 48	213	have been used in the week prior to the first dosing.
49 50	213	<ul> <li>Has not fully recovered from toxicity and/or complications caused by any</li> </ul>
51	214	intervention before starting treatment (i.e., $\leq$ grade 1 or baseline, excluding
52 53	215	weakness or hair loss).
54 55	210	<ul> <li>Known history of human immunodeficiency virus (HIV) infection (i.e. HIV 1/2</li> </ul>
56 57	217	antibody positive).
58 59		
60	219	• Untreated active hepatitis B (defined as HBsAg positive and HBV-DNA copy 7

Page 8 of 17

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

**BMJ** Open

number detected greater than the upper limit of normal value in the laboratory ofthe study center).

- Active HCV-infected subjects (HCV antibody positive and HCV-RNA levels
   above the lower limit of detection).
- Received live vaccine within 30 days prior to the first dose (cycle 1, day 1). Note:
   Injectable inactivated virus vaccine against seasonal influenza is permitted for 30
   days prior to initial administration. However, live attenuated influenza vaccines
   administered intranasally are not permitted.
- Pregnant or lactating women.

Medical history or evidence of disease that may interfere with test results, prevent
 participants from fully participating in the study, abnormal treatment or laboratory
 test values, or other conditions that the investigator considers unsuitable for
 enrollment. The Investigator considers other potential risks unsuitable for
 participation in the study.

234 2.3 Treatment Regimen

Dose selection: tislelizumab: 200mg. pemetrexed: 500mg/m<sup>2</sup>. albumin-bound paclitaxel: 260mg/m<sup>2</sup>. platinum drugs (cisplatin: 60-80mg/m<sup>2</sup>; carboplatin: AUC=5; nedaplatin: 80mg/m<sup>2</sup>). Capsulized FMT: 30 capsules each time. Capsules will be stored in the refrigerator at -80°C. When taking capsules, take them out of the refrigerator, heat them in a water bath (single-hole water bath, JF-01 defroster) at 37°C for 10 minutes, and swallow them with warm water.

Participants will receive FMT combined with tislelizumab + pemetrexed + platinum-based treatment (lung adenocarcinoma) / albumin-bound paclitaxel + platinum-based treatment (lung squamous cell carcinoma) for 4-6 cycles. If there is no progression of the disease after 4-6 cycles of the first-line treatment, then patients will enter the maintenance treatment stage. Patients will receive tislelizumab maintenance treatment (lung squamous cell carcinoma), or tislelizumab + pemetrexed maintenance treatment (lung adenocarcinoma). Treatment continues until disease progression, subject withdraws informed consent, loss of follow-up, or death. 

 Capsulized FMT should be conducted on the 5th day ( $\pm 2$  days) after the start of the first, third, and fifth treatment cycles of chemotherapy, as well as on the 5th day ( $\pm 2$ days) after the first maintenance chemotherapy (lung adenocarcinoma) and before the first maintenance immunotherapy (lung squamous cell carcinoma). All patients will be prohibited from receiving any other treatments with anti-tumor activity or potential anti-tumor activity during the study. This will be monitored by the investigators, the patients, and their families. See figure 2 for details.

**2.4 Study Endpoints** 

The primary endpoint of the study is the 12-month PFS rate. Secondary endpoints include ORR, mPFS, overall survival (OS), duration of response (DoR), safety, microbiome diversity, and quality of life (QoL). Exploratory endpoints including efficacy predictive biomarkers, including fungi, bacteria, metabolomics, and proteomics.

**2.5 Sample Size Estimation** 

The parameters are set as follows: the two-sided significance level alpha is set at 0.05. Based on an estimated dropout rate of 0% and a reference 12-month PFS rate of 35% published in the RATIONALE-304 and RATIONALE-307 clinical studies, it is inferred that the 12-month PFS rate in this trial can reach 55%.<sup>[2,3]</sup> Therefore, a total sample size of 62 participants is determined for this study to ensure statistical power greater than 80%. Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

269 2.6 Efficacy and Safety Assessments270 Imaging evaluation

The assessments of the screening period should be conducted within 28 days prior to the first administration of the study drug. Prior to treatment, researchers at the study center will confirm that subjects have measurable lesions that met the RECIST 1.1 criteria. The methods used to assess tumor burden at baseline must be consistent with those used for each subsequent follow-up assessment (CT/MRI). Additional imaging assessments of other suspected involved areas (e.g., brain) may be conducted based on the subject's clinical symptoms and signs. After enrollment, tumor status will be

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

evaluated using imaging methods every 6 weeks ( $\pm$  7 days), every 12 weeks ( $\pm$  7 days) after 48 weeks until disease progression (RECIST 1.1) or death (during the course of patient treatment). For subjects who have completed treatment or have discontinued treatment for reasons other than disease progression, a single tumor imaging assessment should be performed at the time of treatment completion/discontinuation.

283 Safety evaluation

Researchers will conduct safety assessments on patients every 3 weeks using NCI-CTCAE 5.0. After treatment ends, participants will be followed up for 30 days to detect adverse events (AE). If the patient has not received new anti-tumor treatment within 90 days after the last dose, severe adverse events (SAE) occurring within 90 days after the last dose will be collected. If the subject has received new anti-tumor treatment, SAE prior to the new treatment will be collected, with precedence given to those that have already occurred. Investigators should grade and record adverse events for each subject according to the NCI-CTCAE 5.0 criteria during the study and follow-up period. The characteristics of adverse events will be assessed and recorded based on severity, causality, toxicity grading, management measures, and outcomes.

# 294 Fecal and peripheral blood collection time

With the permission of the ethics committee, patients should provide 10ml whole blood samples and fecal samples at baseline, after two cycles of treatment, before maintenance treatment, and after two cycles of maintenance treatment for the detection of efficacy prediction markers (each cycle is 21 days).

- **2.7 Data Management**
- 300 Data management methods

The study will use a data collection system (91trial) for data management. All subjects will be assigned a unique ID. Researchers will input basic information, drug use, laboratory examination and other raw data information of subjects into 91trial. The system is subject to superior monitoring and cannot be arbitrarily modified. The data manager will write a data audit report based on the trial protocol and audit criteria in the database. The results of the patient's report on tests, examinations, etc. can be

307 obtained from the electronic case report. Quality of life will be assessed using the
 308 EORTC-QLQ-C30 questionnaire, which will be available in both paper and electronic
 309 formats.

Throughout the data collection process, researchers will implement preventive measures to ensure the confidentiality of the documents and safeguard against the identification of participants. All data will be monitored by the data monitoring committee.

-

# **Reporting and collection of adverse events**

When a clinical adverse event occurs, it will be detailed on the case report form with the time of occurrence, clinical manifestations, course of treatment, duration, outcome, and the relationship to this study. For those with laboratory abnormalities, follow-up will continue until the test results return to normal, or to the level before medical intervention, or until it is determined that the event is unrelated to the medical intervention. In the event of a serious adverse event, the serious adverse event form should be filled out and reported to the department and relevant functional departments within 24 hours and recorded in the hospital quality control system (HQS) system, with documentation in the medical record. 

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

**2.8 Statistical Analysis** 

This study will use SPSS statistical software for data analysis. By recording patients' survival times, Kaplan-Meier survival curves for PFS%, PFS and OS will be plotted to visually display the trend of survival rates over time. The occurrence of AE and SAE will be recorded, and the frequency and percentage of different levels of adverse events will be used to describe their occurrence. The Log-rank test will be used to compare whether there are statistically significant differences between the historical control group and the experimental group.

Subgroups will be divided based on patients' demographic characteristics (age, sex),
tobacco use, nutritional status, ECOG performance status, pathological features (tumor
type, solid tumor stage, pathological subtype), metastatic sites (such as brain metastasis,
liver metastasis), PD-L1 expression, and other relevant factors. A multivariate analysis

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

of PFS and OS will be conducted using the Cox proportional hazards model. A p-value
of <0.05 will be considered statistically significant.</li>

### 338 2.9 Trial status

The study was registered at ClinicalTrials.gov (NCT06403111) on May 7, 2024. Enrolment is currently in progress. The first patient was enrolled on June 25, 2024, and the study is expected to end on June 1, 2026.

# **2.10 Patient and public involvement**

343 None.

# **3.** Ethics and Dissemination

The study was approved by the ethics committee of the Second Peoples' Hospital of Changzhou (number [2024] YLJSA005) and is being conducted in accordance with the principles of the Declaration of Helsinki. The study was registered in ClinicalTrials.gov (NCT06403111). Important protocol modifications will be communicated to relevant parties and published on ClinicalTrials.gov. Before participating in the study, the participants (or their legal representatives) will sign the informed consent form. During the study, participants will be provided with any new information that may affect their decision to continue. They can withdraw at any time without facing any penalties or losing any benefits to which they are entitled. The results of this study will be disseminated in a peer-reviewed journal and in conference reports. 

#### **4. Discussion**

The application of ICIs has brought significant benefits to patients with driver genenegative NSCLC, but statistics show that the 5-year survival rate for advanced NSCLC patients receiving immunotherapy is only about 20%.<sup>[13]</sup> With a deeper exploration of the mechanisms of immune resistance, researchers have found that the gut microbiota plays a crucial role in regulating the immune system.

361 Preclinical studies have found that specific gut microbiota or fecal microbiota
 362 transplantation using feces from patients who respond to ICIs can modulate the immune

system, enhance immune cell infiltration in tumors, induce tumor regression, and improve the antitumor efficacy of ICIs.<sup>[8,14–16]</sup> Existing clinical trials have shown that exogenous microflora supplementation by FMT can change the composition of intestinal microflora and the proportion of dominant microflora, reprogram the tumor microenvironment, and potentially reverse immune resistance.<sup>[11,17]</sup> However, the majority of related studies have focused on melanoma, with limited research in the field of lung cancer treatment. Recently, a proof-of-concept clinical trial conducted by a team of researchers in Korea has demonstrated the potential benefits of FMT in a clinical setting other than melanoma, giving us confidence to carry out our study.<sup>[12]</sup> 

Our study is a single-arm clinical trial, which has the limitations of a small sample size and the absence of a control group. So, the conclusions may not be convincing. At the same time, some information derived from a single-arm trial, such as QoL and efficacy predictive biomarkers was relatively limited. Though the results of randomized controlled trials are more convincing, it demands significant investments of human, material, and financial resources.<sup>[18]</sup> On this basis, randomized phase II trials seem to be an effective research method, which requires a relatively small sample size compared to traditional randomized controlled trials, and the inclusion of a control group makes it more scientifically rigorous than a single-arm study.<sup>[19]</sup> Overall, each research method has its own strengths and disadvantages. Therefore, we should consider it carefully in the light of the actual situation. 

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

#### **References**

1. Beay F, Laversanne M, Sung H, et al. Global cancer statistics 2022: GLOBOCAN estimates of
incidence and mortality worldwide for 36 cancers in 185 countries. *CA Cancer J Clin*2024;74(3):229-263 doi:10.3322/caac.21834 [published Online First: 4 April 2024].

2. Lu S, Wang J, Yu Y, et al. Tislelizumab Plus Chemotherapy as First-Line Treatment for Locally
Advanced or Metastatic Nonsquamous NSCLC (RATIONALE 304): A Randomized Phase 3 Trial. *J Thorac Oncol* 2021;16(9):1512-1522 doi:10.1016/j.jtho.2021.05.005 [published Online First: 23
May 2021].

391 3. Wang J, Lu S, Yu X, et al. Tislelizumab plus chemotherapy versus chemotherapy alone as first392 line treatment for advanced squamous non-small-cell lung cancer: final analysis of the randomized,
393 phase III RATIONALE-307 trial. *ESMO Open* 2024, 9(10) doi:10.1016/j.esmoop.2024.103727
394 [published Online First: 25 September 2024].

4. Nagasaki J, Ishino T, Togashi Y. Mechanisms of resistance to immune checkpoint inhibitors.
 396 *Cancer Sci* 2022;113(10):3303-3312 doi:10.1111/cas.15497 [published Online First: 30 July 2022].

1 2 3

397

Enseignement Superieur (ABES) Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

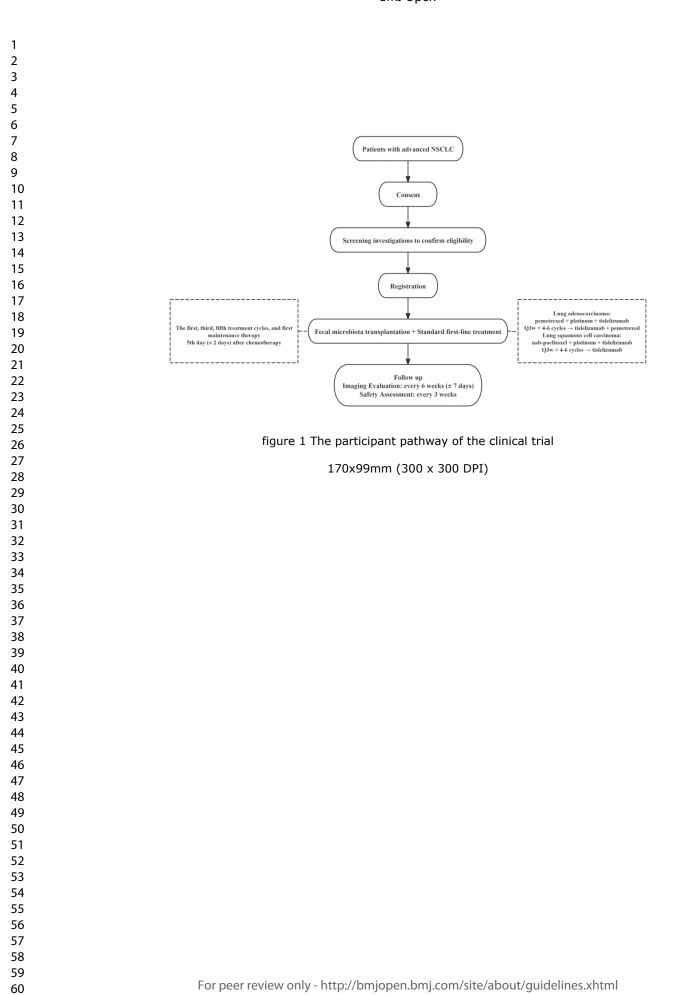
5. Bagchi S, Yuan R, Engleman EG. Immune Checkpoint Inhibitors for the Treatment of Cancer: 4 398 Clinical Impact and Mechanisms of Response and Resistance. Annu Rev Pathol 2021;16:223-249 5 399 doi:10.1146/annurev-pathol-042020-042741 [published Online First: 16 November 2020]. 6 7 400 6. Wu J, Wang S, Zheng B, Qiu X, et al. Modulation of Gut Microbiota to Enhance Effect of 8 401 Checkpoint Inhibitor Immunotherapy. Front Immunol 2021;12:669150 9 402 doi:10.3389/fimmu.2021.669150 [published Online First: 29 June 2021]. 10 11 403 7. Vétizou M, Pitt JM, Daillère R, et al. Anticancer immunotherapy by CTLA-4 blockade relies on 12 404 the gut microbiota. Science 2015;350(6264):1079-1084 doi:10.1126/science.aad1329 [published 13 405 Online First: 5 November 2015]. 14 15 406 8. Sivan A, Corrales L, Hubert N, et al. Commensal Bifidobacterium promotes antitumor immunity 16 407 and facilitates anti-PD-L1 efficacy. Science 2015;350(6264):1084-1089 17 408 doi:10.1126/science.aac4255 [published Online First: 5 November 2015]. 18 19 409 9. Blake SJ, Wolf Y, Boursi B, et al. Role of the microbiota in response to and recovery from cancer 20 410 therapy. Nat Rev Immunol 2024;24(5):308-325 doi:10.1038/s41577-023-00951-0 [published Online 21 411 First: 6 November 2023]. 22 23 412 10. Erdmann J. How gut bacteria could boost cancer treatments. Nature 2022;607(7919):436-439 24 413 doi:10.1038/d41586-022-01959-7 [published Online First: 19 July 2022]. 25 414 11. Davar D, Dzutsev AK, Mcculloch JA, et al. Fecal microbiota transplant overcomes resistance 26 therapy in melanoma patients. Science 2021; 371(6529): 415 anti–PD-1 595-602 to 27 28 doi:10.1126/science.abf3363 [published Online First: 5 February 2021]. 416 29 417 12. Kim Y, Kim G, Kim S, et al. Fecal microbiota transplantation improves anti-PD-1 inhibitor 30 418 efficacy in unresectable or metastatic solid cancers refractory to anti-PD-1 inhibitor. Cell Host 31 32 419 Microbe 2024;32(8):1380-1393.e9 doi:10.1016/j.chom.2024.06.010 [published Online First: 25 33 420 July 2024]. 34 13. Xing Li, Fei Zhou, Chunxia Su. Advances in the research of mechanisms and strategies in 421 35 422 immunotherapy resistance. Chinese Journal of Metastatic Cancer (in Chinese) 2020,03(03):235-36 37 423 240 doi:10.3760/cma.j.cn101548-20200520-00060 [published Online First: 30 September 2020]. 38 424 14. Mager LF, Burkhard R, Pett N, et al. Microbiome-derived inosine modulates response to 39 425 checkpoint inhibitor immunotherapy. Science 2020; 369(6510): 1481-1489 40 41 426 doi:10.1126/science.abc3421 [published Online First: 13 August 2020]. 42 427 15. Routy B, Le Chatelier E, Derosa L, et al. Gut microbiome influences efficacy of PD-1-based 43 immunotherapy 2018; 359(6371):91-97 428 against epithelial tumors. Science 44 429 45 doi:10.1126/science.aan3706 [published Online First: 2 November 2017]. 46 430 16. Gopalakrishnan V, Spencer CN, Nezi L, et al. Gut microbiome modulates response to anti-PD-47 431 1 immunotherapy in melanoma patients. Science 2018; 359(6371): 97-103 48 432 doi:10.1126/science.aan4236 [published Online First: 2 November 2017]. 49 50 433 17. Baruch EN, Youngster I, Ben-Betzalel G, et al. Fecal microbiota transplant promotes response 51 434 in immunotherapy-refractory melanoma patients. Science 2021; 371(6529): 602-609 52 435 doi:10.1126/science.abb5920 [published Online First: 10 December 2020]. 53 54 436 18. Hariton E, Locascio JJ. Randomised controlled trials - the gold standard for effectiveness 55 437 research: Study design: randomised controlled trials. BJOG: an international journal of obstetrics 56 438 and gynaecology 2018;125(13):1716 doi:10.1111/1471-0528.15199 [published Online First: 19 57 439 June 2018]. 58 59 440 19. Rubinstein LV, Korn EL, Freidlin B, et al. Design issues of randomized phase II trials and a 60

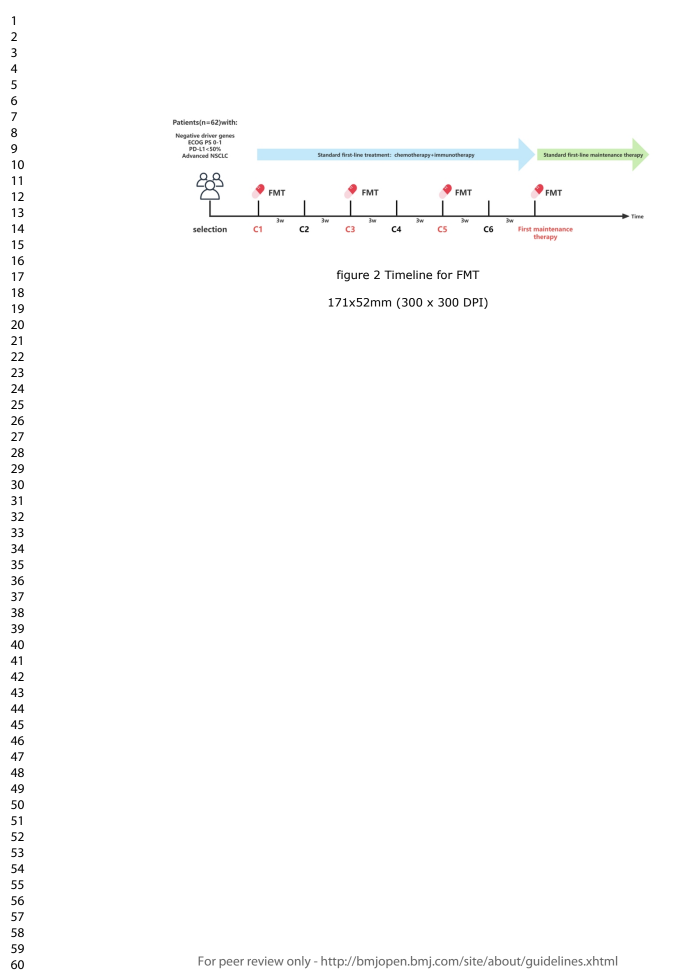
#### **BMJ** Open

proposal for phase II screening trials. <i>J Clin Oncol</i> 2005;23(28):7199-7206 doi:10.1200/JCO.2005.01.149 [published Online First: 1 October 2005].	
Authors' Contributions	
YW and LQ are responsible for the article's concept and writing. QG, DL, and DQ are	
responsible for clinical treatment. QG, LQ, YW, and XW are responsible for data	
collection and compilation. HJ and QG coordinate all the work. YW and LQ contributed	
equally to this paper. QG is the guarantor, reviews the entire study design and the draft.	Prot
All authors review and approve the final manuscript.	tectec
Funding	Enseignement Superieur (AB
This study was supported by the 2022 Clinical Research project of Changzhou Medical	right,
Center, Nanjing Medical University (CMCC202201), 2022 Changzhou 8th Batch of	inclu
Science and Technology Project (Applied Basic Research) (CJ20220086).	iding f
Acknowledgements	or uses
We would like to express our sincere appreciation to all the participants.	eignen relate
Peer review	to text
Not externally peer reviewed.	and da
Data sharing	ata mir
No additional data are available.	ES) . hining, Al training, and similar technologies
Competing interests statement	rainir
There are no associations with commercial entities, nor any financial relationships	lg, an
involving their spouse or children under 18 years of age. Additionally, there are no non-	d sim
financial associations that could be relevant to the submitted manuscript.	ilar tec
Figure legends	hnolog
figure 1 The participant pathway of the clinical trial	ies.
figure 2 Timeline for FMT	
15	

Page 16 of 17

**BMJ** Open





#### Fecal microbiota transplantation combined with platinumbased doublet chemotherapy and Tislelizumab as first-line treatment for driver-gene negative advanced non-small-cell lung cancer (NSCLC): study protocol for a prospective, multi-center, single-arm exploratory trial

Journal:	BMJ Open
Manuscript ID	bmjopen-2024-094366.R2
Article Type:	Protocol
Date Submitted by the Author:	27-Jan-2025
Complete List of Authors:	Wei, Yanshuang; Changzhou Second People's Hospital of Nanjing Medical University, Department of Oncology Qin, Lanqun; Changzhou Second People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Wu, Xinyu; Changzhou Second People's Hospital of Nanjing Medical University, Department of Oncology Li, Dongqing; Changzhou Second People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Qian, Danping; Changzhou Second People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Qian, Danping; Changzhou Second People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center Jiang, Hua; Changzhou Second People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical center Geng, Qian; Changzhou Second People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical center Geng, Qian; Changzhou Second People's Hospital of Nanjing Medical University, Department of Oncology; Nanjing Medical University, Changzhou Medical Center
<b>Primary Subject Heading</b> :	Oncology
Secondary Subject Heading:	Research methods
Keywords:	CHEMOTHERAPY, Clinical Relevance, Clinical Trial, ONCOLOGY, Respiratory tract tumours < ONCOLOGY



1	Fecal microbiota transplantation combined with platinum-based doublet
2	chemotherapy and Tislelizumab as first-line treatment for driver-gene negative
3	advanced non-small cell lung cancer (NSCLC): study protocol for a prospective,
4	multi-center, single-arm exploratory trial
5	Yanshuang Wei <sup>1</sup> †, Lanqun Qin <sup>1,2</sup> †, Xinyu Wu <sup>1</sup> , Dongqing Li <sup>1,2</sup> , Danping Qian <sup>1,2</sup> ,
6	Hua Jiang <sup>1,2*</sup> , Qian Geng <sup>1,2*</sup>
7	(†These authors contributed equally to this work and share first authorship)
8	<sup>1</sup> Department of Oncology, the Second People's Hospital of Changzhou, the Third
9	Affiliated Hospital of Nanjing Medical University, Changzhou, 213000, China.
10	<sup>2</sup> Changzhou Medical Center, Nanjing Medical University, Changzhou, 213003, China.
11	*Correspondence:
12	(Qian Geng is the main correspondence author and Hua Jiang is the secondary
13	correspondence author)
14	Qian Geng, E-mail: karengq@njmu.edu.cn, Address: Department of Oncology, the Second People's
15	Hospital of Changzhou, the Third Affiliated Hospital of Nanjing Medical University, Changzhou,
16	China.
17	Hua Jiang, E-mail: czeyjh@njmu.edu.cn, Address: Department of Oncology, the Second People's
18	Hospital of Changzhou, the Third Affiliated Hospital of Nanjing Medical University, Changzhou,
19	China.
20	ABSTRACT
21	Introduction: Chemotherapy combined with immunotherapy is the standard first-line
22	treatment for driver-gene negative advanced non-small cell lung cancer (NSCLC).
23	However, due to the immune microenvironment imbalance and immune status
24	impairment caused by repeated chemotherapy, as well as the primary or secondary
25	resistance to immune checkpoint inhibitors (ICIs), the efficacy of immunotherapy
26	combined with chemotherapy remains unsatisfactory. Recent studies have shown that
27	fecal microbiota transplantation (FMT) can modulate the intestinal microflora,
28	influence the tumor immune microenvironment, and even enhance the efficacy of
29	immunotherapy. Therefore, we conduct this study to evaluate the efficacy and safety of
30	FMT combined with standard first-line treatment for driver gene-negative advanced
31	NSCLC.
32	Methods and analysis: FMT-JSNO-02 (NCT06403111) is a prospective, multi-center,
33	single-arm exploratory study. It is planned to include 62 cases of previously untreated
34	driver-gene negative, Eastern Cooperative Oncology Group Performance Status
75	$\sim$

35 (ECOG PS) 0-1, Programmed death ligand 1 (PD-L1) <50% advanced NSCLC</li>
 36 patients, who will be given FMT by orally ingested stool capsules on the basis of

standard first-line treatment of chemotherapy combined with immunotherapy. The
safety and efficacy of this treatment regimen will be evaluated, with the primary

<sup>39</sup> endpoint being the 12-months progression-free survival (PFS) rate.

Ethics and dissemination: The study was approved by the ethics committee of the
Second People's Hospital of Changzhou (number [2024] YLJSA005) and is being
conducted in accordance with the principles of the Declaration of Helsinki. The results
of the study will be submitted for publication in peer-reviewed journals.

44 Trial registration number: ClinicalTrials.gov Identifier: NCT06403111. Date of
 45 registration: May 7, 2024, the first version protocol.

#### 46 Strengths and limitations of this study:

- This is a multi-center prospective study.
- Patients receive FMT via orally ingested stool capsules.
- 49 The sample size in this study is relatively small.
- This is a single-arm study rather than a randomized controlled study.

51 keywords: Fecal microbiota transplantation, Non-Small Cell Lung Cancer,
52 Chemotherapy, Immunotherapy

54 1. Introduction

### 55 Current Situation and Dilemmas of Immunotherapy for Non-Small Cell lung 56 cancer (NSCLC)

Lung cancer is the most common malignancy worldwide and remains the leading cause of cancer-related mortality, accounting for 18.7% of all cancer deaths.<sup>[1]</sup>As the main pathological subtype of lung cancer, the majority of NSCLC patients present with local spread or distant metastasis upon diagnosis, losing the opportunity for surgery and having a very poor prognosis. In recent years, immune checkpoint inhibitors (ICIs), particularly those targeting the programmed death receptor 1 (PD-1) and its ligand (PD-L1), markedly improved the treatment outcomes and becoming an effective treatment strategy for NSCLC. 

At present, the combination of chemotherapy and immunotherapy has become the standard first-line treatment for patients with advanced NSCLC due to extensive clinical studies conducted on pembrolizumab and nivolumab. Tislelizumab developed in China has also shown remarkable efficacy. In patients with lung adenocarcinoma, the addition of tislelizumab to chemotherapy resulted in significantly prolonged median Page 3 of 18

#### **BMJ** Open

progression-free survival (mPFS) compared to chemotherapy alone (9.7 versus 7.6 month), with a 12-month PFS rate of 31.3%.<sup>[2]</sup> Similarly, in patients with lung squamous cell carcinoma, the mPFS for tislelizumab plus chemotherapy group was also higher than that of chemotherapy alone (9.6 months versus 5.5 months).<sup>[3]</sup>

Although ICIs significantly improved the survival of patients with advanced NSCLC, primary or acquired immune resistance inevitably occurs in clinical practice. Existing research suggests that primary immune resistance is related to the dysfunction in tumor antigen processing and presentation, insufficient T cell infiltration in the tumor microenvironment (TME), and over expression of suppressive immune cells.<sup>[4]</sup> Acquired immune resistances is related to adaptive changes that occur in tumor cells and the TME during exposure to immunotherapy.<sup>[5]</sup>

Nowadays, a growing body of research suggests that gut flora may be able to reverse immune resistance. For instance, gut microbiota can regulate the infiltration of immune cells in the TME by modulating innate and adaptive immunity. They can also reprogram the TME through their reactive metabolites or secretions, affecting the efficacy of ICIs and even reversing immune resistance.<sup>[6]</sup> Researchers have conducted preliminary explorations into how gut microbiota affects the efficacy of immunotherapy. Zitvoge found that the antitumor effects of cytotoxic T-lymphocyte-associated protein 4 (CTLA-4) blockade depended on distinct Bacteroides species. Tumors in antibiotic-treated or germ-free mice did not respond to CTLA-4 blockade. This defect was overcome by gavage with B. fragilis, by immunization with B. fragilis polysaccharides, or by adoptive transfer of B. fragilis-specific T cells.<sup>[7]</sup> At the same time, Sivan et al. found that mice orally administered Bifidobacterium could achieve the same anti-tumor effect as PD-L1 monoclonal antibodies, and the combination almost completely inhibited tumor growth.<sup>[8]</sup> Therefore, modulating gut microbes has the potential to further improve the treatment outcomes for patients receiving immunotherapy. 

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

100 Current Status of Fecal Microbiota Transplantation (FMT)

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

> The gut microbiota is susceptible to various factors, including the use of broad-spectrum antibiotics, repeated chemotherapy and other treatments. Common methods to regulate the intestinal microbiota include dietary intervention, probiotics, prebiotics, and FMT. Among these approaches, FMT stands as the most established technique.<sup>[9]</sup> FMT is the transplantation of functional flora from the stool of a healthy person into the intestine of a patient in a certain way. This method mainly includes two ways: injecting microflora fluids and swallowing capsules. Among them, FMT by orally ingested stool capsules (capsulized FMT) has the characteristics of a wide range of application, high acceptance degree, relative economy and convenience.<sup>[10]</sup>

The relationship between intestinal flora and immunotherapy efficacy has been preliminarily confirmed in some clinical studies of malignant tumors. For example, in a study focused on 15 melanoma patients resistant to anti-PD-1 therapy, rechallenge with FMT and Pembrolizumab showed objective responses in 3 out of 15 patients, and these 3 patients kept stable disease (SD) for more than 12 months.<sup>[11]</sup> In a clinical trial (NCT04264975), researchers performed FMT with anti-PD-1 inhibitors in 13 patients with anti-PD-1 refractory advanced solid cancers. The results showed that FMT induced sustained microbiota changes and clinical benefits in 6 patients, achieving an objective response rate (ORR) of 7.7% and a disease control rate (DCR) of 46.2%.<sup>[12]</sup> As a result, FMT holds great potential to reverse immune resistance and enhance the efficacy of immunotherapy. 

Hence, we conduct such a prospective, exploratory study in order to evaluate the efficacy and safety of FMT combined with standard first-line treatment for driver genenegative advanced NSCLC. We expect that the addition of FMT to the standard firstline treatment in NSCLC will be safe and will have an enhancing effect.

- **2. Methods**
- 128 2.1 Study Design

129 This is a prospective, multi-center, single-arm exploratory study aimed at evaluating

Page 5 of 18

#### **BMJ** Open

the efficacy and safety of FMT combined with standard first-line treatment for driver-gene negative NSCLC patients. The trial protocol was written in accordance with the Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT) guidance. The study will mainly be conducted at the Second People's Hospital of Changzhou, the Third Affiliated Hospital of Nanjing Medical University. Recruitment information can be obtained through outpatient consultations, posters displayed in the wards, and other channels. A total of 62 patients with driver-gene negative, Eastern Cooperative Oncology Group (ECOG) performance status (PS) 0-1, and PD-L1<50% who have not received prior treatment are expected to be included in this study without grouping. Patients will receive platinum-based doublet chemotherapy combined with tislelizumab as standard first-line treatment for 4-6 cycles and then enter maintenance therapy. FMT will be given to patients during the first, third, and fifth treatment cycles, as well as during the first maintenance therapy. All the enrolled patients or their designated agents will sign the informed consent form within 3 days before starting treatment. This study was approved by the ethics committee of the Second People's Hospital of Changzhou (number [2024] YLJSA005). See figure 1 for details.

## 146 2.2 Patient Population147 Inclusion criteria:

The subjects voluntarily joined the study and were able to sign the informed
 consent with good compliance.

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

• Age 18-80 years old (when signing the informed consent form).

 Patients with histologically or cytologically proven locally advanced (IIIB/IIIC), metastatic, or recurrent (stage IV) NSCLC who are inoperable and unable to receive radical concurrent chemoradiotherapy, according to the International Association for the Study of Lung Cancer and the American Joint Committee on Cancer Classification, 8th Edition TNM Classification of Lung cancer.

- Have not received systemic intravenous anti-tumor therapy before, and the driver gene is negative.
- 158 PD-L1 expression < 50%.
- According to the solid tumor efficacy evaluation criteria (RECIST version 1.1),

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

there is at least one radiographically measurable lesion. That is, in CT or MRI detection, the longest diameter of a single lesion was  $\geq$  10mm, or the pathological enlargement of a single lymph node was  $\geq$  15mm.

• The physical status score of Eastern Tumor Collaboration Group (ECOG) was 0-

1.

• Expected survival > 3 months.

Have adequate organ and bone marrow function, laboratory tests within 7 days prior to enrollment meet the following requirements (no blood components, cell growth factors, albumin or other corrective drugs are allowed within 14 days prior to obtaining laboratory examination), as follows: 1) Blood routine: absolute neutrophil count (ANC)  $\geq 1.5 \times 10^{9}$ /L, platelet (PLT)  $\geq 75 \times 10^{9}$ /L, hemoglobin  $(HGB) \ge 90$  g/L (no blood transfusion or erythropoietin dependence within 14 days). 2) Liver function: serum total bilirubin (TBIL)  $\leq 2$  times the upper limit of normal (ULN). Alanine aminotransferase (ALT) and/or aspartate aminotransferase  $(AST) < 5 \times ULN$ , serum albumin > 28 g/L, alkaline phosphatase  $(ALP) < 5 \times ULN$ . 3) Renal function: serum creatinine (Cr)  $\leq 1.5 \times ULN$ , or creatinine clearance  $\geq 50$ mL/min (using the standard Cockcroft-Gault formula): Urine routine results showed urinary protein < 2+. For patients with urine protein  $\ge 2+$  at baseline, 24-hour urine collection and 24-hour urine protein quantification < 1g should be performed. 4) Coagulation function: International standardized ratio (INR) or prothrombin time (PT)  $\leq$  1.5 times ULN. If the subject is receiving anticoagulant therapy, as long as the INR is within the intended range of anticoagulant drug use. For female subjects of reproductive age, a urine or serum pregnancy test should be performed and the result is negative 3 days prior to receiving the initial study drug administration. 

Subjects and their sexual partners are required to use a medically approved contraceptive method (such as an intrauterine device (IUD), contraceptive pill, or condom) during the study treatment period and for 6 months after the end of the study treatment period.

1 2		
3 4	189 Exclusion criteria:	
5 6	190	• Currently participating in an interventional clinical study or receiving another
7 8	191	investigational drug or investigational device within 4 weeks prior to initial dosing.
9	192	• Received proprietary Chinese medicines with anti-tumor indications or
10 11	193	immunomodulatory drugs (thymosin, interferon, interleukin, etc.) within 2 weeks
12 13	194	before the first administration, or received major surgical treatment within 3 weeks
14 15	195	before the first administration.
16 17	196	• Class III - IV congestive heart failure (New York Heart Association classification),
18 19	197	poorly controlled and clinically significant arrhythmias.
20 21	198	• Any arterial thrombosis, embolism or ischemia, such as myocardial infarction,
22 23	199	unstable angina pectoris, cerebrovascular accident or transient ischemic attack,
24 25	200	occurred within 6 months before treatment.
26 27	201	• Known allergic reaction to the drug in this study.
28 29	202	• Patients who require long-term oral, intravenous, or intramuscular administration
30 31	203	of systemic corticosteroids.
32	204	• Symptomatic central nervous metastases. Patients with asymptomatic brain
33 34	205	metastases (BMS) or BMS whose symptoms are stable after treatment are eligible
35 36	206	to participate in this study if they meet all of the following criteria: measurable
37 38	207	lesions outside the central nervous system. No midbrain, pontine, cerebellum,
39 40	208	meninges, medulla oblongata or spinal cord metastasis. Maintain clinical stability
41 42	209	for at least 2 weeks. Stop hormone therapy 3 days before the first dose of the study
43 44	210	drug.
45 46	211	• There is an active infection requiring treatment or systemic anti-infective drugs
47 48	212	have been used in the week prior to the first dosing.
49 50	213	• Has not fully recovered from toxicity and/or complications caused by any
50 51 52	214	intervention before starting treatment (i.e., $\leq$ grade 1 or baseline, excluding
52 53 54	215	weakness or hair loss).
55	216	<ul> <li>Known history of human immunodeficiency virus (HIV) infection (i.e. HIV 1/2</li> </ul>
56 57	210	antibody positive).
58 59	217	<ul> <li>Untreated active hepatitis B (defined as HBsAg positive and HBV-DNA copy</li> </ul>
60	210	<ul> <li>Ontreated active hepatitis D (defined as fibsAg positive and fibv-DNA copy</li> <li>7</li> </ul>

Page 8 of 18

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

**BMJ** Open

number detected greater than the upper limit of normal value in the laboratory ofthe study center).

Active HCV-infected subjects (HCV antibody positive and HCV-RNA levels
 above the lower limit of detection).

Received live vaccine within 30 days prior to the first dose (cycle 1, day 1). Note:
 Injectable inactivated virus vaccine against seasonal influenza is permitted for 30
 days prior to initial administration. However, live attenuated influenza vaccines
 administered intranasally are not permitted.

• Pregnant or lactating women.

Medical history or evidence of disease that may interfere with test results, prevent
 participants from fully participating in the study, abnormal treatment or laboratory
 test values, or other conditions that the investigator considers unsuitable for
 enrollment. The Investigator considers other potential risks unsuitable for
 participation in the study.

**2.3 Treatment Regimen** 

Dose selection: tislelizumab: 200mg. pemetrexed: 500mg/m<sup>2</sup>. albumin-bound paclitaxel: 260mg/m<sup>2</sup>. platinum drugs (cisplatin: 60-80mg/m<sup>2</sup>; carboplatin: AUC=5; nedaplatin: 80mg/m<sup>2</sup>). Capsulized FMT: 30 capsules each time. Capsules will be stored in the refrigerator at -80°C. When taking capsules, take them out of the refrigerator, heat them in a water bath (single-hole water bath, JF-01 defroster) at 37°C for 10 minutes, and swallow them with warm water.

Participants will receive FMT combined with tislelizumab + pemetrexed + platinum-based treatment (lung adenocarcinoma) / albumin-bound paclitaxel + platinum-based treatment (lung squamous cell carcinoma) for 4-6 cycles. If there is no progression of the disease after 4-6 cycles of the first-line treatment, then patients will enter the maintenance treatment stage. Patients will receive tislelizumab maintenance treatment (lung squamous cell carcinoma), or tislelizumab + pemetrexed maintenance treatment (lung adenocarcinoma). Treatment continues until disease progression, subject withdraws informed consent, loss of follow-up, or death. 

 Capsulized FMT should be conducted on the 5th day ( $\pm 2$  days) after the start of the first, third, and fifth treatment cycles of chemotherapy, as well as on the 5th day ( $\pm 2$ days) after the first maintenance chemotherapy (lung adenocarcinoma) and before the first maintenance immunotherapy (lung squamous cell carcinoma). All patients will be prohibited from receiving any other treatments with anti-tumor activity or potential anti-tumor activity during the study. This will be monitored by the investigators, the patients, and their families. See figure 2 for details.

**2.4 Study Endpoints** 

The primary endpoint of the study is the 12-month PFS rate. Secondary endpoints include ORR, mPFS, overall survival (OS), duration of response (DoR), safety, microbiome diversity, and quality of life (QoL). Exploratory endpoints including efficacy predictive biomarkers, including fungi, bacteria, metabolomics, and proteomics.

**2.5 Sample Size Estimation** 

The parameters are set as follows: the two-sided significance level alpha is set at 0.05. Based on an estimated dropout rate of 0% and a reference 12-month PFS rate of 35% published in the RATIONALE-304 and RATIONALE-307 clinical studies, it is inferred that the 12-month PFS rate in this trial can reach 55%.<sup>[2,3]</sup> Therefore, a total sample size of 62 participants is determined for this study to ensure statistical power greater than 80%. Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

268 2.6 Efficacy and Safety Assessments269 Imaging evaluation

The assessments of the screening period should be conducted within 28 days prior to the first administration of the study drug. Prior to treatment, researchers at the study center will confirm that subjects have measurable lesions that met the RECIST version 1.1 criteria. The methods used to assess tumor burden at baseline must be consistent with those used for each subsequent follow-up assessment (CT/MRI). Additional imaging assessments of other suspected involved areas (e.g., brain) may be conducted based on the subjects' clinical symptoms and signs. After enrollment, tumor status will

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

be evaluated using imaging methods every 6 weeks ( $\pm$  7 days), every 12 weeks ( $\pm$  7 days) after 48 weeks until disease progression (RECIST version 1.1) or death (during the course of patient treatment). For subjects who have completed treatment or have discontinued treatment for reasons other than disease progression, a single tumor should performed imaging assessment be at the time of treatment completion/discontinuation.

283 Safety evaluation

Researchers will conduct safety assessments on patients every 3 weeks using NCI-CTCAE 5.0. After treatment ends, participants will be followed up for 30 days to detect adverse events (AE). If the patient has not received new anti-tumor treatment within 90 days after the last dose, severe adverse events (SAE) occurring within 90 days after the last dose will be collected. If the subject has received new anti-tumor treatment, SAE prior to the new treatment will be collected, with precedence given to those that have already occurred. Investigators should grade and record adverse events for each subject according to the NCI-CTCAE 5.0 criteria during the study and follow-up period. The characteristics of adverse events will be assessed and recorded based on severity, causality, toxicity grading, management measures, and outcomes.

#### 294 Fecal and peripheral blood collection time

With the permission of the ethics committee, patients should provide 10ml whole blood samples and fecal samples at baseline, after two cycles of treatment, before maintenance treatment, and after two cycles of maintenance treatment for the detection of efficacy prediction markers (each cycle is 21 days).

**2.7 Data Management** 

#### **300 Data management methods**

The study will use a data collection system (91trial) for data management. All subjects will be assigned a unique ID. Researchers will input basic information, drug use, laboratory examination and other raw data information of subjects into 91trial. The system is subject to superior monitoring and cannot be arbitrarily modified. The data manager will write a data audit report based on the trial protocol and audit criteria in

the database. The results of the patients' report on tests, examinations, etc. can be
obtained from the electronic case report. Quality of life will be assessed using the
EORTC-QLQ-C30 questionnaire, which will be available in both paper and electronic
formats.

Throughout the data collection process, researchers will implement preventive measures to ensure the confidentiality of the documents and safeguard against the identification of participants. All data will be monitored by the data monitoring committee.

**Reporting and collection of adverse events** 

When a clinical adverse event occurs, it will be detailed on the case report form with the time of occurrence, clinical manifestations, course of treatment, duration, outcome, and the relationship to this study. For those with laboratory abnormalities, follow-up will continue until the test results return to normal, or to the level before medical intervention, or until it is determined that the event is unrelated to the medical intervention. In the event of a serious adverse event, the serious adverse event form should be filled out and reported to the department and relevant functional departments within 24 hours and recorded in the hospital quality control system (HQS) system, with documentation in the medical record. 

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

**2.8 Statistical Analysis** 

This study will use SPSS statistical software for data analysis. By recording patients' survival times, Kaplan-Meier survival curves for PFS%, PFS and OS will be plotted to visually display the trend of survival rates over time. The occurrence of AE and SAE will be recorded, and the frequency and percentage of different levels of adverse events will be used to describe their occurrence. The Log-rank test will be used to compare whether there are statistically significant differences between the historical control group and the experimental group.

Subgroups will be divided based on patients' demographic characteristics (age, sex),
tobacco use, nutritional status, ECOG performance status, pathological features (tumor
type, solid tumor stage, pathological subtype), metastatic sites (such as brain metastasis,

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

liver metastasis), PD-L1 expression, and other relevant factors. A multivariate analysis
of PFS and OS will be conducted using the Cox proportional hazards model. A p-value
of <0.05 will be considered statistically significant.</li>

#### 338 2.9 Trial status

The study was registered at ClinicalTrials.gov (NCT06403111) on May 7, 2024. Enrolment is currently in progress. The first patient was enrolled on June 25, 2024, and the study is expected to end on June 1, 2026.

- **2.10 Patient and public involvement**
- 343 None.

 **3. Ethics and Dissemination** 

The study was approved by the ethics committee of the Second People's Hospital of Changzhou (number [2024] YLJSA005) and is being conducted in accordance with the principles of the Declaration of Helsinki. The study was registered in ClinicalTrials.gov (NCT06403111). Important protocol modifications will be communicated to relevant parties and published on ClinicalTrials.gov. Before participating in the study, the participants (or their legal representatives) will sign the informed consent form. During the study, participants will be provided with any new information that may affect their decision to continue. They can withdraw at any time without facing any penalties or losing any benefits to which they are entitled. If participants suffer harm due to research interventions or research-related procedures during the trial, appropriate compensation will be provided in accordance with relevant laws, regulations, and the guidance of the ethics committee. The results of this study will be disseminated in a peer-reviewed journal and in conference reports. 

#### **4. Discussion**

The application of ICIs has brought significant benefits to patients with driver genenegative NSCLC, but statistics show that the 5-year survival rate for advanced NSCLC patients receiving immunotherapy is only about 20%.<sup>[13]</sup> With a deeper exploration of

Page 13 of 18

#### **BMJ** Open

the mechanisms of immune resistance, researchers have found that the gut microbiotaplays a crucial role in regulating the immune system.

Preclinical studies have found that specific gut microbiota or fecal microbiota transplantation using feces from patients who respond to ICIs can modulate the immune system, enhance immune cell infiltration in tumors, induce tumor regression, and improve the antitumor efficacy of ICIs.<sup>[8,14–16]</sup> Existing clinical trials have shown that exogenous microflora supplementation by FMT can change the composition of intestinal microflora and the proportion of dominant microflora, reprogram the tumor microenvironment, and potentially reverse immune resistance.<sup>[11,17]</sup> However, the majority of related studies have focused on melanoma, with limited research in the field of lung cancer treatment. Recently, a proof-of-concept clinical trial conducted by a team of researchers in Korea has demonstrated the potential benefits of FMT in a clinical setting other than melanoma, giving us confidence to carry out our study.<sup>[12]</sup> 

Our study is a single-arm clinical trial, which has the limitations of a small sample size and the absence of a control group. So, the conclusions may not be convincing. At the same time, some information derived from a single-arm trial, such as QoL and efficacy predictive biomarkers was relatively limited. Though the results of randomized controlled trials are more convincing, it demands significant investments of human, material, and financial resources.<sup>[18]</sup> On this basis, randomized phase II trials seem to be an effective research method, which requires a relatively small sample size compared to traditional randomized controlled trials, and the inclusion of a control group makes it more scientifically rigorous than a single-arm study.<sup>[19]</sup> Overall, each research method has its own strengths and disadvantages. Therefore, we should consider it carefully in the light of the actual situation. 

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies.

#### **References**

1. Beay F, Laversanne M, Sung H, et al. Global cancer statistics 2022: GLOBOCAN estimates of
incidence and mortality worldwide for 36 cancers in 185 countries. *CA Cancer J Clin*2024;74(3):229-263 doi:10.3322/caac.21834 [published Online First: 4 April 2024].

2. Lu S, Wang J, Yu Y, et al. Tislelizumab Plus Chemotherapy as First-Line Treatment for Locally
 Advanced or Metastatic Nonsquamous NSCLC (RATIONALE 304): A Randomized Phase 3 Trial.

*J Thorac Oncol* 2021;16(9):1512-1522 doi:10.1016/j.jtho.2021.05.005 [published Online First: 23 May 2021]. 1 2 3

4

5

6

7

8

9

10 11

12

13

14

15

16

17

18 19

20

21

22

23 24

25

26

27 28

29

30

31 32

33

34

35

36 37

38

39

40 41

42

43

44

45 46

47

48

49 50

51

52

53 54

55

56

3. Wang J, Lu S, Yu X, et al. Tislelizumab plus chemotherapy versus chemotherapy alone as first-394 395 line treatment for advanced squamous non-small-cell lung cancer; final analysis of the randomized, phase III RATIONALE-307 trial. ESMO Open 2024, 9(10) doi:10.1016/j.esmoop.2024.103727 396 [published Online First: 25 September 2024]. 397 398 4. Nagasaki J, Ishino T, Togashi Y. Mechanisms of resistance to immune checkpoint inhibitors. 399 Cancer Sci 2022;113(10):3303-3312 doi:10.1111/cas.15497 [published Online First: 30 July 2022]. 5. Bagchi S, Yuan R, Engleman EG. Immune Checkpoint Inhibitors for the Treatment of Cancer: 400 401 Clinical Impact and Mechanisms of Response and Resistance. Annu Rev Pathol 2021;16:223-249 402 doi:10.1146/annurev-pathol-042020-042741 [published Online First: 16 November 2020]. 403 6. Wu J, Wang S, Zheng B, Qiu X, et al. Modulation of Gut Microbiota to Enhance Effect of 404 Checkpoint Inhibitor Immunotherapy. Front Immunol 2021;12:669150 405 doi:10.3389/fimmu.2021.669150 [published Online First: 29 June 2021]. 406 7. Vétizou M, Pitt JM, Daillère R, et al. Anticancer immunotherapy by CTLA-4 blockade relies on 407 the gut microbiota. Science 2015;350(6264):1079-1084 doi:10.1126/science.aad1329 [published 408 Online First: 5 November 2015]. 409 8. Sivan A, Corrales L, Hubert N, et al. Commensal Bifidobacterium promotes antitumor immunity 410 anti-PD-L1 efficacy. Science 2015;350(6264):1084-1089 and facilitates 411 doi:10.1126/science.aac4255 [published Online First: 5 November 2015]. 9. Blake SJ, Wolf Y, Boursi B, et al. Role of the microbiota in response to and recovery from cancer 412 therapy. Nat Rev Immunol 2024;24(5):308-325 doi:10.1038/s41577-023-00951-0 [published Online 413 414 First: 6 November 2023]. 10. Erdmann J. How gut bacteria could boost cancer treatments. Nature 2022;607(7919):436-439 415 416 doi:10.1038/d41586-022-01959-7 [published Online First: 19 July 2022]. 417 11. Davar D, Dzutsev AK, Mcculloch JA, et al. Fecal microbiota transplant overcomes resistance to anti-PD-1 therapy in melanoma patients. Science 2021; 371(6529): 595-602 418 419 doi:10.1126/science.abf3363 [published Online First: 5 February 2021]. 420 12. Kim Y, Kim G, Kim S, et al. Fecal microbiota transplantation improves anti-PD-1 inhibitor efficacy in unresectable or metastatic solid cancers refractory to anti-PD-1 inhibitor. Cell Host 421 422 Microbe 2024;32(8):1380-1393.e9 doi:10.1016/j.chom.2024.06.010 [published Online First: 25 423 July 2024]. 424 13. Xing Li, Fei Zhou, Chunxia Su. Advances in the research of mechanisms and strategies in 425 immunotherapy resistance. Chinese Journal of Metastatic Cancer (in Chinese) 2020,03(03):235-426 240 doi:10.3760/cma.j.cn101548-20200520-00060 [published Online First: 30 September 2020]. 427 14. Mager LF, Burkhard R, Pett N, et al. Microbiome-derived inosine modulates response to 428 checkpoint inhibitor immunotherapy. Science 2020: 369(6510): 1481-1489 429 doi:10.1126/science.abc3421 [published Online First: 13 August 2020]. 430 15. Routy B, Le Chatelier E, Derosa L, et al. Gut microbiome influences efficacy of PD-1-based 431 immunotherapy against epithelial tumors. Science 2018; 359(6371):91-97 432 doi:10.1126/science.aan3706 [published Online First: 2 November 2017]. 433 16. Gopalakrishnan V, Spencer CN, Nezi L, et al. Gut microbiome modulates response to anti-PD-434 1 immunotherapy in melanoma patients. Science 2018; 359(6371): 97-103 435 doi:10.1126/science.aan4236 [published Online First: 2 November 2017]. 14

436 17. Baruch EN, Youngster I, Ben-Betzalel G, et al. Fecal microbiota transplant promotes response
437 in immunotherapy-refractory melanoma patients. *Science* 2021; 371(6529): 602-609
438 doi:10.1126/science.abb5920 [published Online First: 10 December 2020].

439 18. Hariton E, Locascio JJ. Randomised controlled trials - the gold standard for effectiveness
440 research: Study design: randomised controlled trials. *BJOG: an international journal of obstetrics*441 *and gynaecology* 2018;125(13):1716 doi:10.1111/1471-0528.15199 [published Online First: 19
442 June 2018].

443 19. Rubinstein LV, Korn EL, Freidlin B, et al. Design issues of randomized phase II trials and a
444 proposal for phase II screening trials. *J Clin Oncol* 2005;23(28):7199-7206
445 doi:10.1200/JCO.2005.01.149 [published Online First: 1 October 2005].

446 Authors' Contributions

447 YW and LQ are responsible for the article's concept and writing. QG, DL, and DQ are
448 responsible for clinical treatment. QG, LQ, YW, and XW are responsible for data

- 449 collection and compilation. HJ and QG coordinate all the work. YW and LQ contributed
- 450 equally to this paper. QG is the guarantor, reviews the entire study design and the draft.
- 451 All authors review and approve the final manuscript.

#### 452 Funding

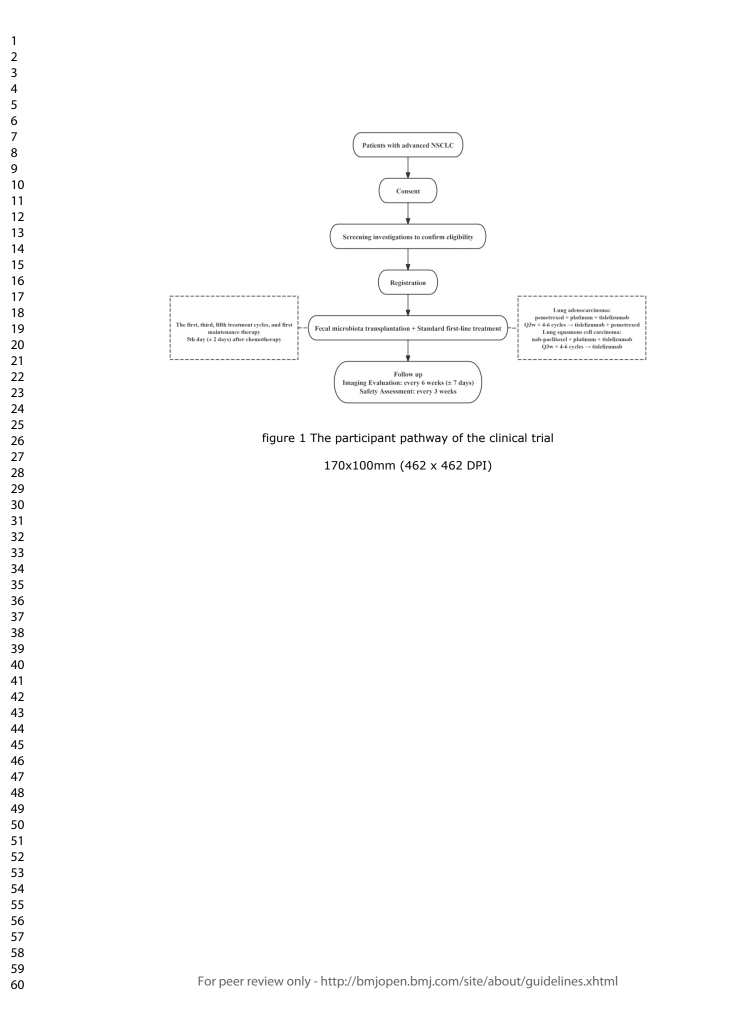
453 This study was supported by the 2022 Clinical Research project of Changzhou Medical

- 454 Center, Nanjing Medical University (CMCC202201), 2022 Changzhou 8th Batch of
- 455 Science and Technology Project (Applied Basic Research) (CJ20220086).
  - 456 Acknowledgements
  - 457 We would like to express our sincere appreciation to all the participants.
  - 458 Peer review
  - 459 Not externally peer reviewed.
  - 460 Data sharing
  - 461 No additional data are available.
  - **Competing interests**
  - 463 Not applicable.
- 464 Figure legends
- **figure 1** The participant pathway of the clinical trial

Enseignement Superieur (ABES) . Protected by copyright, including for uses related to text and data mining, Al training, and similar technologies.

**figure 2** Timeline for FMT

to beet teriewony



BMJ Open: first published as 10.1136/bmjopen-2024-094366 on 4 March 2025. Downloaded from http://bmjopen.bmj.com/ on June 12, 2025 at Agence Bibliographique de l Enseignement Superieur (ABES)

