

New Publication: Mistletoe Extract in Patients with Advanced Pancreatic Cancer

The randomised, placebo-controlled, double-blind MISTRAL trial on mistletoe therapy in pancreatic cancer has now been published internationally by the German Medical Journal/Deutsches Ärzteblatt (*Dtsch Arztebl Int* 2024; 121: 347–54). The trial was initiated and coordinated by the first author, Dr. Kathrin Wode, designed in collaboration with German experts on mistletoe research, Swedish scientists and clinicians, and funded by research foundations.

The MISTRAL trial investigated whether mistletoe therapy can further improve the results of optimal, modern oncological palliative treatment. The study explored whether mistletoe extracts, given alongside comprehensive oncological treatment and palliative care, can further prolong overall survival and improve health-related quality of life in patients with pancreatic cancer. In addition to mistletoe therapy, participants in the MISTRAL trial received optimal palliative medical care according to today's standards (including chemotherapy in 80% of cases).

In the MAPAC study from 2013/14, the main intervention was mistletoe therapy – with the result of a significant improvement in quality of life and overall survival. In the MISTRAL study, concomitant mistletoe therapy did not show any additional advantage in terms of survival time and quality of life. Frequent concerns about the safety and interaction profile of mistletoe therapy with chemotherapy could be ruled out. Further data on aspects of quality of life, the development of body weight as a measure of general health, biomarkers (laboratory values) and qualitative measures of subjective well-being are still being analysed.

What was the MISTRAL trial design?

Since patients with advanced pancreatic cancer have a short survival time and only limited treatment options available, the double-blind, randomised, placebo-controlled MISTRAL trial investigated whether mistletoe extracts (ME), given in addition to comprehensive oncological treatment and palliative care, can prolong overall survival (OS) and improve health-related quality of life (HRQoL) in 290 patients in 9 Swedish oncology centres. The main inclusion criteria were advanced, exocrine pancreatic cancer and an ECOG performance status between 0 and 2. Participants were randomly assigned to either the mistletoe therapy group (n = 143) or the placebo group (n = 147), stratified according to trial centre and recommendation of palliative chemotherapy. ME or placebo was injected subcutaneously three times a week for nine months and HRQoL was measured seven times during this period. Participants were also given medication for symptom control (analgesics, antiemetics, anxiolytics, glucocorticoids) and, if necessary, multidisciplinary palliative medical care, delivered at home or at the hospital).

What were the results?

There was no statistically significant advantage for mistletoe therapy either on the OS, or the HRQoL dimensions “global health” and “quality of life”. The number, severity and outcome of adverse events were also comparable between the mistletoe and placebo groups, except for the more frequent local skin reactions under ME (66% vs. 1%). The adjusted hazard ratio for OS was 1.13 [0.89; 1.44], the median survival time was 7.8 months in the ME group and 8.3 months in the placebo group, and the first look at HRQoL (EORTC-QLQ C30 questionnaire, questions 29 and 30: ‘How would you rate your overall state of health during the last week’ and ‘How would you rate your overall quality of life during the last week’) was also comparable in both groups ($p = 0.86$).

Has MISTRAL refuted the MAPAC trial which recorded significant improvements in quality of life and lifespan in patients with advanced pancreatic cancer due to mistletoe therapy?

No, because the treatments given alongside mistletoe therapy differ considerably between the two trials: The MAPAC study (1, 2), published in 2013/14, investigated the effect of mistletoe therapy as the main intervention in a group of patients with advanced pancreatic cancer. In these cases, palliative chemotherapy was no longer considered for these patients in the situation at the time. Concomitant treatment in Serbia consisted of medications to hold symptoms like pain, nausea/vomiting and dyspepsia in check.

The participants in the MISTRAL study received mistletoe therapy in addition to optimal palliative medical care by today's standards. As an expression of the difference between the two studies, the survival time of patients without mistletoe therapy was 2.7 months in the MAPAC study compared to 8.3 months in the MISTRAL study. In the MAPAC trial, the survival time of patients with mistletoe therapy was increased to 4.8 months, while the survival time of 7.9 months for mistletoe patients in the MISTRAL trial did not differ significantly from the group without mistletoe therapy. According to the publication, the modern supportive treatments may have caused a statistical “ceiling effect”, leaving not much to be improved by an additional mistletoe therapy.

The MAPAC trial was controlled and randomised, but not blinded and not placebo-controlled. Must the conclusion of the MISTRAL trial therefore be deemed more credible?

Mistletoe therapy poses a particular challenge for blinding because it usually leads to a characteristic local skin reaction at the injection site. The authors of the MAPAC study decided against blinding for fear of having to discontinue the trial if patients’ belonging to the treatment group became revealed.

The MISTRAL trial publication states that 93 out of 140 patients who received mistletoe therapy reported a local skin reaction and therefore unblinding may have occurred in 65% of patients in the observation group. However, the data shows this had no influence on the results.

Do these findings conclude the research into mistletoe therapy for pancreatic cancer?

No, as the complete data from the MISTRAL trial has not yet been conclusively analysed. In addition to survival time and global quality of life, other aspects of quality of life, the development of body weight as a measure of general health, biomarkers (laboratory values)

and a qualitative interview series with the patients were also documented. All this data has yet to be analysed and published.

Were there any positive results from the MISTRAL trial?

Yes; due to a lack of data, in the past, there were some concerns about the safety and interaction profile of mistletoe therapy with chemotherapy. The MISTRAL trial has now answered both questions. According to the authors, there were no reasons for safety concerns: only two cases of severe side effects (1 urticaria (“hives”) and 1 pseudo-allergic reaction) occurred, which was in line with the known side effect profile of mistletoe therapy; the frequency of less than 1.5% occurrence was also in line with expectations based on the existing scientific literature.

Furthermore, the researchers found “no evidence of an influence of ME (mistletoe extracts) on the effect of chemotherapy in terms of prolonged or shortened survival or toxicity. However, this is only an assessment, as our study was not designed to investigate this question. However, this assessment is supported by a phase I study (3) which found no effect of ME on the metabolism of gemcitabine - the most commonly used chemotherapy for pancreatic cancer.”

What does this mean?

Following the MAPAC trial from 2013/14, in which the question on mistletoe’s efficacy was answered positively, the Swedish researchers wanted to investigate whether mistletoe therapy could be used to further improve the results of optimal, modern oncological palliative treatment in a disease where modern medicine has reached its limits. Comprehensive palliative medical treatment can significantly increase the survival time of pancreatic cancer patients (4), but within this specific setting mistletoe therapy statistically is not likely to offer any additional benefits.

The patients in the MISTRAL study already had a much better survival time (8.3 months) due to the integrated palliative medical treatment provided in Sweden, than the patients in the MAPAC trial from 2013/14, who only received reduced symptomatic therapy to control pain and nausea (2.7 months). From the MISTRAL study, it may be concluded that mistletoe therapy does not provide any additional benefit when comprehensive modern palliative oncological care is provided for pancreatic cancer patients. This may be different within different therapeutic settings. Therefore, as recommended in the German S3 guideline ‘Complementary medicine in oncology’, the decision as to whether to choose concomitant mistletoe therapy for any given pancreatic cancer situation should remain with the patients and their treating specialists, taking into account their individual situation and specific needs.

When is further data from the MISTRAL trial expected to be published?

According to the authors, further publications are currently being prepared. However, they are obliged to maintain confidentiality about the exact timeline. Iscador AG as the manufacturer of the study medication will be informed immediately prior to each subsequent publication by the authors in a conference. It remains our topmost priority to process the new results promptly and to make them available in summarised form on our homepage.

Original publication

- The publication is openly accessible [in English](#) and [in German](#)
Wode K, Kienle GS, Björ O, et al. Mistletoe extract in patients with advanced pancreatic cancer: a double-blind, randomized, placebo-controlled trial (MISTRAL). *Dtsch Arztebl Int* 2024; 121: 347–54. DOI: 10.3238/arztebl.m2024.0080
- A short interview with the trial's main investigator, Dr Kathrin Wode (in German):
<https://www.aerzteblatt.de/archiv/239602/3-Fragen-an-Kathrin-Wode>

Further Literature

- (1) Tröger W, Galun D, Reif M, et al. *Viscum album* [L.] extract therapy in patients with locally advanced or metastatic pancreatic cancer: a randomised clinical trial on overall survival. *Eur J Cancer* 2013;49:3788-3797. DOI: [10.1016/j.ejca.2013.06.043](https://doi.org/10.1016/j.ejca.2013.06.043)
- (2) Tröger W, Galun D, Reif M, et al. Quality of life of patients with advanced pancreatic cancer during treatment with mistletoe: a randomized controlled trial. *Dtsch Arztebl Int* 2014;111:493-502. DOI: [10.3238/arztebl.2014.0493](https://doi.org/10.3238/arztebl.2014.0493)
- (3) Mansky PJ, Wallerstedt DB, Sannes TS, et al. NCCAM/NCI phase 1 study of mistletoe extract and gemcitabine in patients with advanced solid tumors. *Evid Based Complement Alternat Med* 2013; ID964592. DOI: 10.1155/2013/964592
- (4) Seufferlein T, Porzner M, Heinemann V, Tannapfel A, Stuschke M, Uhl W: Duktales Pankreaskarzinom: Chirurgische Therapie, pathologische Aufarbeitung des Präparats, neoadjuvante, adjuvante und palliative Therapie. *Dtsch Arztebl International* 2014; 111(22):396-402. DOI: 10.3238/arztebl.2014.0396