# Mistletoe Extracts in Cancer Care

# **Healthcare Provider Resource**

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## **General information**

#### Proper name:

Viscum album Loranthaecea, Viscum album L.

#### Common names:

Mistletoe, European Mistletoe, Viscum album extracts (VAE)

#### Routes of administration:

Subcutaneous (SC), intravenous (IV), intramuscular, intrapleural, intratumoral, and intravesical instillation. This monograph will focus on the two most common routes: SC and IV.

#### Commercially available products:

Helixor®, Iscador®, abnobaVISCUM® (Isorel®, Lektinol®, Eurixor® are no longer available)

<u>Common uses in cancer care:</u> Mistletoe extracts are commonly used to enhance immune function, support quality of life, reduce cancer-related side effects and symptoms, slow disease progression, reduce risk of recurrence, and improve survival.

# Summary

Viscum album extracts (VAE) are used in integrative cancer care to support immune function, reduce side effects, improve quality of life (QOL), and possibly improve survival and recurrence. The most common routes of administration are subcutaneous (SC) injection and intravenous (IV) infusion; most research pertains to SC administration. Proposed mechanisms of action include immunomodulation of both innate and adaptive immune response, and direct cytotoxicity. Increased lymphocytes (T cells, B cells, and NK cells), dendritic cells, cytokines including INF-gamma and IL6, and presence of IgG antibodies to mistletoe lectins and viscotoxins have been observed. SC and IV VAE are well tolerated; serious side effects such as allergy and anaphylaxis are rare but have been reported. Mild and self-limiting side effects including local injection site reactions (with SC use), fatigue, and mild fever are common. Studies in people with cancer have found that mistletoe is likely to support QOL, reduce symptom burden, and reduce side effects associated with treatment when given alongside standard care. Studies on survival and tumor response are not conclusive; some studies find benefit and others find no difference compared to control groups. VAE is not a cancer cure and not an alternative to conventional care. Overall methodological quality is poor, and studies with better methodology are less likely to find benefit to survival. In conclusion, mistletoe is a promising adjunctive therapy for QOL and side effect management, but more research is needed from well controlled studies to further elucidate its impact on survival and recurrence risk for people with cancer.

### **Background**

Preparations from European Mistletoe are used as complementary treatment for people with cancer, most notably in Germany (1). Mistletoe, a parasitic plant from the Santalacea family, is commonly prepared as an extract and is commercially available from several manufacturers. The extracts contain various compounds which vary slightly based on host tree, harvest time and preparation method. Available products are often named based on host tree, commonly including malus (apple tree: "M"), abies (fir tree: "A"), pinus (pine: "P"), and quercus (oak: "Qu") (1, 2). Some mistletoe extracts are fermented (Iscador®), while others are unfermented (Helixor®, abnobaVISCUM®).

This monograph discusses evidence pertaining to the use of European mistletoe (*Viscum album* L) extracts in complementary cancer care, omitting American and Korean mistletoe, and pharmaceutical preparations (e.g., E. coli-derived recombinant counterpart of mistletoe lectin-I known as rViscumin (Aviscumine)) (3, 4). This monograph primarily discusses the subcutaneous and intravenous routes of administration, which are most often used in North America. Throughout this summary, mistletoe will be referred to as VAE (*Viscum album* extract) or mistletoe.

## Methods

Monographs are created by the Patterson Institute for Integrative Oncology research team and are updated approximately every two years. Comprehensive and structured literature searches were performed in Medline and Cochrane library from inception for English-language studies in people with cancer. Additional scoping reviews were performed by research staff to obtain supporting information such as background information, mechanism of action, and safety data. Articles are duplicate-screened, data is extracted into standardized spreadsheets, and studies summarized using descriptive statistics.

### **Pharmacokinetics**

Pharmacokinetic data on VAE is limited. A phase I study evaluated the pharmacokinetics of VAE by administering a single SC injection of abnobaVISCUM Fraxini (20mg) to 15 healthy male volunteers (5). Mistletoe lectins were detected in all serum samples after injection, with mean and median peak concentrations reached 1 and 2 hours after injection, respectively. Concentration-time profiles considerably, indicating non-linear kinetics, and thus half-life could not be determined (5). Mistletoe lectins were detectable in 60% of the men after 14 days. Significant individual variability in subcutaneous mistletoe pharmacokinetics exists. Pharmacokinetics of other VAE administration routes have not been studied. In vitro research has found no cytochrome P450 induction capacity of VAE, and no inhibition over 50% when concentrations equivalent to 100,000 times the clinically relevant dose in plasma were used. Thus, the authors concluded that herb-drug interactions due to P450 interactions were unlikely (6).

# **Mechanism of Action**

Active compounds of VAE include mistletoe lectins (ML) (I, II and III), viscotoxin (VT) proteins, flavonoids, phenylpropanoids, triterpenes, phytosterol,

alkaloids, polyalcohols, and polysaccharides (7). Lectins and viscotoxins have been studied the most (2, 8). Different VAE formulas contain varied concentrations of MLs and VTs due to host tree, time of harvest, and extraction method, and thus the biological response is also expected to differ (2). The two primary mechanisms of action for VAE are immune system modulation and cytotoxicity.

#### Immunologic activity:

Lectins are proposed to be primarily responsible for the immunologic activity of VAE (9). While diverse effects have been noted, overall, most studies report immune function improvement with VAE administration (2). Immune parameters observed to increase or improve include granulocytes (neutrophils, eosinophils, basophils), lymphocytes (T cells, B cells, NK cells), dendritic cells, cytokines and interleukins (including IFN-g, TNF-a, IL-1, IL-4, IL-5, IL-6), and IgG antibodies (2, 10-12).

Randomized trials in healthy volunteers indicate that SC VAE stimulates both innate and adaptive immune responses (9, 13, 14). One study randomized 43 healthy volunteers to SC VAE, purified mistletoe lectin (ML), ML-free VAE, or placebo twice weekly for 8 weeks, and analyzed differential blood counts and peripheral blood mononuclear cells (PBMC) (9). Significant increases in leukocyte, granulocyte, and antigen-induced production of GM-CSF, IL-5, and IFN gamma by PBMC with VAE and ML treatment compared to placebo groups was observed. Another study compared SC injections of Iscucin Populi (IP), Visum Mali (VM), or placebo and demonstrated eosinophilia with both VAEs, increased CD4 T-lymphocytes in the VAE IP group, and no change in IL6 or CRP in any group (13). An adaptive immune response to VAE was demonstrated in a 12week trial of 47 people randomized to Iscador Q (rich in ML), Iscador P (rich in viscotoxins, low in ML), or placebo (14). Anti-ML-1 IgG antibodies were present in all Iscador Q-treated subjects but only 6 exposed to Iscador P. Anti-VA2 IgG-antibodies were detected in all individuals in VAE groups, none of the participants receiving placebo developed antibodies.

Studies in cancer populations report similar results. A small RCT of women with breast cancer receiving adjuvant chemo-radiotherapy found that 7 weeks of VAE significantly increased IFN-g and IL-6 compared to control (15). In a study of 98 women with breast cancer having surgery, a single infusion of 1mg Iscador M one-hour prior to anaesthetic prevented the surgical suppression of granulocyte function when compared to the control group (16). However, results of four controlled trials of VAE during adjuvant chemotherapy for breast (n= 3) and gastric (n=1) cancer found that VAE did not improve neutrophil counts (the most abundant granulocyte) as there was no change compared to controls (17-20). Details of these four studies can be found in Table 1.

Natural killer (NK) cells are of particular interest in cancer research. Two studies have found improvements in NK cell numbers or function in people treated with VAE peri-operatively. One RCT randomized 70 people undergoing surgery for digestive tract cancer to receive VAE for 4 weeks peri-operatively or control (21). The treatment had significantly group less immunosuppressive effects from surgery compared to controls, with an increased number of lymphocytes including NK cells, T cells and B cells, and an increase in immunoglobulins. A study of patients undergoing surgery for colon cancer found similar results, showing that perioperative infusion of VAE prevented NK suppression 24h post-surgery in the mistletoe group (22).

Lastly, VAE may exert effects on dendritic cells (DCs). VAE stimulates both the maturation and the activity of the DCs and counteracts the immunosuppressive effect of tumour cells on DCs as evident from in vitro and in vivo studies (10-12). Several other studies presented in tables 1-3 provide additional information on the immune effects of VAE administration.

#### Cytotoxic activity:

Mistletoe lectins, viscotoxins and alkaloids are believed to be responsible for mistletoe's cytotoxic activity (23). Proposed mechanisms include protein synthesis inhibition, triggering apoptosis and necrosis, indirect cytotoxic effects resulting from cytokine release, and increasing natural killer cell cytotoxicity and macrophage activity (23-25). Most studies on the cytotoxic activity of VAE come from preclinical data. It has been suggested that although low doses of VAE have been effective for supporting immune function, higher doses may be needed to exert cytotoxic effects which may also increase toxicity and side effects of the therapy (23).

#### Other actions

Mistletoe may attenuate markers of inflammation, which may result in improved fatigue, as demonstrated by one study in women with early-stage breast cancer (26).

# Clinical Evidence Related to Effectiveness

There are 14 clinical trials (18 publications) for SC VAE in cancer (Table 1), 2 clinical trials for IV VAE in cancer (Table 2), 7 studies using other routes of VAE administration (Table 3), and 24 observational studies (Table 4) identified from the literature search. These studies are discussed below based on administration route and outcomes assessed. The most up to date systematic reviews and meta-analyses are also discussed, as in many cases they contain data from studies not meeting our inclusion criteria (e.g. Germanlanguage, or journals not indexed by Medline or Cochrane), and thus provide additional information.

# **Subcutaneous injections**

There are a diverse number of human studies using SC VAE injections, though they vary in quality and design. There are 14 clinical trials described in 18 publications (Table 1), as well as several observational studies (Table 4). Overall, VAE appears to likely benefit immune function, QOL, and reduce disease- and treatment-related symptoms. Results are mixed regarding tumour

response and survival. Variance in survival studies may be attributed to differences in VAE preparations, dosing, cancer types, administration schedules and study design. Several systematic reviews report methodological concerns within published clinical trials (8, 27-31).

#### **Ouality of Life**

Of the 14 subcutaneous VAE clinical trials identified, 12 investigated endpoints related to QOL, side-effects and/or toxicity of cancer treatments (17-21, 32-38). Eleven were randomised controlled trials (17-21, 32-35, 37, 38), only one of which was placebo-controlled (35). Five studies included patients with breast cancer (17, 18, 20, 35, 37), four studied patients with pancreatic cancer (21, 33, 38, 39), two each with colorectal cancer (21, 36), lung cancer (32, 37), and gastric cancer (19, 21), and one each with relapsed osteosarcoma (34), esophageal cancer (21) and ovarian cancer (37).

The majority of studies report that VAE improves QOL endpoints observed across different cancer types, conventional treatments, and stages of disease. Only one study reported that VAE did not improve QOL but did reduce treatment related toxicity (32). Most studies report mixed QOL benefit, with some endpoints significantly improving while others not. While VAE appears to consistently improve aspects of QOL, predictions of which *specific* endpoints will be improved vary between patients. Due to methodological issues and trial heterogeneity, the exact type and magnitude of benefit warrants further investigation.

Nine studies used the same validated standardized QOL assessment tool (EORTC QLQ-C30) (17-20, 32-34, 40, 41), allowing for inter-study QOL endpoint comparison. VAE significantly improved global health in relapsed osteosarcoma patients (34), gastric cancer patients receiving chemotherapy (19), advanced pancreatic cancer patients receiving supportive care (41), breast cancer patients receiving chemotherapy (17, 18, 20, 40, 42), but no benefit for patients with lung cancer receiving carboplatin chemotherapy (32). Two studies reported that VAE application resulted in significant benefit for physical functioning (20, 38). VAE

significantly benefited role functioning in four studies, three of which included patients with breast cancer receiving chemotherapy (17, 18, 20) and one which evaluated patients with advanced pancreatic cancer (41). Five studies observed significant benefit of VAE application regarding emotional functioning, including three with breast cancer patients receiving chemotherapy (17, 18, 20), one with relapsed osteosarcoma patients post-surgery (34), and one in advanced pancreatic cancer receiving best supportive care (38). Lastly, social and cognitive function were significantly improved compared to control patients in a study of patients with advanced pancreatic cancer (38).

Ten studies reported use of VAE during different chemotherapy treatments (17-20, 32-34, 37, 40, 42), of which only one reported that no significant benefit was noted for QOL (32). Chemotherapy agents included carboplatin based treatments (32), cyclophosphamide, doxorubicin plus 5-Fluorouracil (5FU), (17, 18, 20), cyclophosphamide, methotrexate, and 5-FU (40, 42), doxifluridine (5-DFUR) (19), and "mixed/multiple" types (37).

The most recent systematic review and meta-analysis of VAE for QOL in patients with cancer was published in 2020 (43). In this review, 26 prospective controlled trials with two or more arms were included and comprised 30 data sets (25 RCTs, 5 CTs). Compared to control groups, the post-treatment standardized mean difference in global QoL was d=0.61 (95% CI 0.41-0.81, p<0.00001), indicating a medium-sized, clinically meaningful effect favoring mistletoe. Studies included various types of cancer, conventional treatments, and applied various brands of subcutaneous mistletoe preparations. There was a high risk of bias due to lack of blinding and heterogeneity across studies. Other systematic reviews show similar results (1, 8, 29, 30, 44, 45), with one exception which concluded no benefit from mistletoe (28).

#### Symptom management and treatment toxicity

It is likely that at least part of the documented improvements in QOL is attributable to the effects of mistletoe on managing symptoms and toxicities, particularly in relation to chemotherapy (37, 46). Evidence from a range of study designs suggests a benefit for VAE treatment in symptom management and chemotherapy toxicity. Side effects and toxicities which may be improved include nausea, vomiting, diarrhea, appetite loss, pain, fatigue, weight loss, non-hematological toxicities in general, and need for chemotherapy dose-reductions. Further research from high quality studies is needed, as methodological quality continues to be a concern.

A randomized controlled study of patients with stage III and IV lung cancer receiving carboplatin-based chemotherapy found that VAE decreased the frequency of chemotherapy dose reductions (44% vs 13%, p=0.005), grade 3-4 non-hematological toxicities (41% vs 16%, p=0.043) and hospitalisations (54% vs 24%, p=0.016) (32). No benefit was found for hematological toxicities (grade 3-4). An open label study of patients with metastatic treatment-resistant colorectal cancer initiating VAE reported that 40% of participants experienced symptomatic relief of nausea, vomiting, diarrhea, constipation, fatigue and dyspnea (36). One RCT administering VAE during 5-DFUR to patients with early-stage gastric cancer reported a significantly lower rate of diarrhea in the intervention group compared to control (p=0.014) (19).

Several specific symptoms have been improved with the use of VAE in clinical trials. Pain scores significantly improved in five studies (published in 6 reports) (17, 18, 20, 34, 39, 41) and failed to improve in three (47-49), all of which used the EORTC QLQ-C30 for QOL assessment. Appetite loss significantly improved in four studies (17, 18, 20, 41). Fatigue scores significantly improved with VAE use in three-clinical-studies (20, 34, 41) and in one observational study (50), possibly by attenuating markers of inflammation (26). Finally, insomnia and weight loss improved with the use of VAE compared to a control group in patients with advanced pancreatic cancer (41), in this study weight increased by 5.3% in the VAE arm compared to a 3.2 % weight loss in the control arm.

The 2020 systematic review discussed previously (43) included a meta-analysis on QOL subdomains including

specific symptoms across 10 studies. The standardized mean difference (SMD) of VAE compared to control in seven of 14 QOL dimensions were statistically significant in favor of mistletoe (p<0.05). Although all symptoms improved with VAE, only nausea and vomiting, pain, dyspnea and diarrhea met statistical significance (fatigue, insomnia, appetite loss and constipation did not). One systematic review included seven studies which specifically assessed chemotherapy-related side effects. Five of seven studies documented significant benefit with VAE (30). Another systematic review published in German included 10 studies that assessed mistletoe in combination with chemotherapy (51) and documents inconsistent results ranging from no effect to positive effects. Other systematic reviews have found similar findings regarding chemotherapy toxicity (28).

#### Survival and tumor response

Six of the clinical trials described in table 1 investigated survival and/or tumor response endpoints in different cancer populations (18, 32-34, 36, 52). The studies evaluated patients with lung cancer (32, 52), breast cancer (18, 52), pancreatic cancer (33, 52), colorectal cancer (36, 52) and relapsed osteosarcoma (34, 53). Several observational studies and systematic reviews have also been published and are briefly described.

From English-language clinical trials (Table 1), survival outcomes are mixed, with two trials and a long-term follow-up on one reporting a survival benefit (33, 34, 53), two reporting no effect (18, 32) and two studies having no comparator to determine effect (36, 52). Several systematic reviews and meta-analyses of mistletoe for survival have been published; all reporting that some, but not all studies, show a survival benefit (1, 27, 30, 31, 44, 45, 54-56). Notably, methodological quality is a concern, and studies with better methodologies were less likely to find a significant benefit.

The two studies showing a significant survival benefit investigated patients with advanced pancreatic cancer (33) and relapsed osteosarcoma (34), which published

long-term follow up results in 2020 (53). In a phase III RCT, 220 patients with stage III or IV pancreatic cancer, receiving standard supportive care were randomized to VAE or control. Median overall survival was 4.8 and 2.7 months in the VAE and control groups, respectively (p <0.0001) (33). An RCT of 20 patients with relapsed osteosarcoma (stages I-III) randomized participants to VAE or etoposide after surgery (34). Post-relapse disease free survival (PRDFS) at 1 year was 55.6% in the VAE group compared to 12% in historical controls, and 27.3% in the etoposide group. Median PRDFS was 39 months (2-73 months) in the VAE group and 4 months (1-47 months) in the etoposide group (34). A 2020 follow-up on this RCT assessed PRDFS 144 months later. The median PRDFS was 106 months and 7 months, in the VAE and etoposide groups, respectively. The 10-year overall survival (OS) rates were estimated to be 64% in the Viscum arm and 33% in the etoposide arm (53).

The two studies that did not show a survival benefit from the use of mistletoe included a study of patients with stage III and IV non-small-cell lung cancer receiving carboplatin based chemotherapy (32) and a study in patients with non-metastatic breast cancer receiving surgery and adjuvant chemotherapy (18).

Several observational studies have reported benefit with VAE. A retrospective observational study of 240 patients with advanced stage pancreatic cancer compared survival time for those receiving VAE therapy and those not. The study found that the combination of VAE and chemotherapy significantly improved survival compared to chemotherapy alone (12.1 vs 7.3 months, p=0.014). In patients not receiving chemotherapy (supportive care only), patients receiving VAE lived significantly longer (5.4 vs 2.5 months, p=0.006) (57). A retrospective study of 158 patients with stage IV NSCLC, primarily receiving subcutaneous VAE, reported that compared to chemotherapy alone, those receiving concomitant VAE had a significantly better median survival (17 months compared to 8 months) (p=0.007) (58). A retrospective cohort study looked at the use of SC VAE alongside neoadjuvant chemoradiotherapy pre-operatively, in patients with stage II-III rectal adenocarcinoma (59). In the mistletoe group (n = 15) compared to the control group (n = 37) there were significantly better outcomes for pathologic complete response rate (53.5% vs 21.6%, p=0.044), tumor regression grade (66.7% vs 32.4%, p=0.024), T downstaging (86.7% vs 43.2%, p=0.004), overall TNM downstaging (86.7% vs 56.8%, p=0.040), and presence of lymphovascular invasion (13.3% vs 32.4%, p=0.04).

The most recent systematic review and meta-analysis (2020) of 32 controlled trials (13, 745 patients) reported on overall and event-free survival from studies on Iscador published from 1963-2014 (31). The overall survival hazard ratio (HR) was 0.59 (CI: 0.53 to 0.65, p<0.0001), favouring Iscador treatment. None of the studies were blinded, and funnel plot analysis found a moderate performance bias, thus, results should be interpreted with caution. On subgroup analysis, hazard ratios for survival were statistically significantly in favor of Iscardor in breast, cervical, colorectal, liver metastases, uterine, ovarian, pancreatic, and stomach cancer, and not significantly improved in lung, osteosarcoma, or skin cancer. The most recent systematic review to evaluate all types of subcutaneous mistletoe was published in 2019 (27). Fourteen randomized controlled trials were included, and 5/14 studies found significant benefit for survival in breast cancer, advanced stage glioma, non-metastatic uterine cancer and pancreatic cancer. Nine studies found no overall survival benefit in patients with breast cancer, colorectal cancer, gynecological cancer, lung cancer and melanoma. Most studies found no significant effect for progression free survival, disease specific survival or disease-free survival. Similar to the 2020 review, study extensively, methodology varied with heterogeneity observed between trials for cancer type, stage of disease, VAE administration, concomitant treatments and survival measures. While most studies ranked low for reporting bias, major methodological concerns including selection bias, performance bias, attrition bias and the issue of multiple testing were identified in most studies.

In addition to the above-mentioned data, there are many care reports and case series that have been published. These are not reported in this monograph given the availability of higher quality evidence. However, in areas where research is limited (as in subsequent sections), case reports have been included given the paucity of data.

In summary, while both positive and neutral data exists, due to inter-study heterogeneity and methodological issues, no conclusive statement can be made regarding the benefit of VAE for cancer survival. However, the research on mistletoe for survival outcomes in pancreatic cancer (33, 57) and osteosarcoma (34, 53) is compelling. More research is needed.

#### **Intravenous infusion**

Two clinical trials investigated the effects of intravenous VAE administration; one phase I study primarily pertaining to safety (60) and one RCT evaluating survival (61) (Table 2). The phase 1 clinical study investigated escalating doses (200mg to 2000mg) of VAE in people with varied advanced cancers, but no concurrent cancer treatment. There were no serious AEs related to the IV VAE. The authors report that 2/21 patients had an unexpected positive clinical response observed by tumor marker changes and 1/21 had slowed progression (60). The study reporting on survival was a 3-arm RCT of 64 patients with advanced colorectal cancer comparing adjuvant chemotherapy, adjuvant chemotherapy + VAE, and surgery without adjuvant treatment (61). Median survival in the adjuvant VAE group was significantly longer (757 days) compared to both the chemotherapy alone group (545 days, p < 0.05) and the surgery alone group (502 days, p< 0.05). There were fewer side effects in the VAE group compared to chemotherapy alone group (0% vs 19%).

#### Mixed routes of administration

Four observational studies and one systematic review with meta-analysis combined data on patients administered VAE using different routes of administration, commonly SC, IV, and intratumoral. Of the observational studies, two included NSCLC patients, one included pancreatic cancer patients, and the fourth looked at patients with breast cancer (62-65). The pancreatic and NSCLC studies used mistletoe (either

SC, IV, intratumoral or combined) plus standard oncologic treatment, and found survival outcomes favoring the combined approach which were also costeffective compared to standard oncologic treatment alone (62, 63). The second study among NSCLC patients yielded non-significant overall survival benefits, however, subgroup analysis revealed that patients with unresected tumours were more likely to benefit (64). A longitudinal study on patients with breast cancer analyzed the impact of SC and IV VAE on cancer related fatigue and QOL. Participants were analyzed based on four groups: those receiving VAE only, chemotherapy, chemotherapy and VAE, or no chemotherapy or VAE (could receive endocrine or immunotherapy). (65). Patients receiving VAE without chemotherapy experienced significant improvements on thermo-coherence (an aspect of internal homeostasis related to subjective comfort in body temperature), fatigue, and seven EORTC subscales at 24 months. Chemo-, immuno- and endocrine therapies resulted in declines in fatigue scores by 6-17 points, whereas the VAE group improved 12 points. Similarly, the VAE group improved in insomnia and physical functioning scores while these scores worsened in conventional care groups. However, these results should be interpreted with caution due to the methodology of this study, and given VAE use alone may not be a great comparator to chemotherapy.

Three case reports described outcomes for patients treated with both IV and SC mistletoe. Two cases showed\_long-term disease-free survival in patients with stage IV renal cell carcinoma. In one, VAE was used alongside chemoimmunotherapy (66), and in the other VAE was applied as monotherapy (67). The third case report was in a patient with dedifferentiated high-grade liposarcoma in the retroperitoneum who survived 10.5 years with good QOL with conventional treatments in addition to IV and SC VAE (68).

# Other routes of administration (excluding IV and SC)

VAE has been applied through other routes aside from subcutaneous and intravenous administration including:

intravesicular, intratumoral, intrapleural and intraperitoneal applications. The related research is not described in this monograph; however, details for these alternate routes are listed in Table 3. Case reports exist, but are not reviewed in this monograph.

# **Applications with limited research**

## Hematological malignancies

Two case reports and one observational study were identified for VAE in hematological malignancies. One case report describes a 65-year-old male with diffuse Bcell lymphoma who received R-CHOP chemotherapy, initially experiencing a minor response. The addition of VAE to chemotherapy, and then continuation of application afterwards resulted in further regression, with the patient in complete remission at time of publication (69). A second case report on two patients with primary cutaneous B-cell lymphoma describes regression of disease (no conventional treatment provided) with the combined use of high dose IV, subcutaneous and intra-tumoral VAE administration (70). Authors report that both patients were in remission 3.5 years after commencement of VAE treatment. A German language retrospective observational study reported that patients with a hematological malignancy (types not specified) who received VAE (n=205) had a median survival of 11.4 years compared to 8.6 years from the controls used (n=9), these results were not statistically significant (71). There were no cases where mistletoe was associated with deterioration.

Data is limited regarding safety and efficacy of VAE use for hematological cancers at this time.

#### Pediatric use

Two retrospective studies were identified related to pediatric application of mistletoe. One was a retrospective case series of ten children with varied relapsed or advanced cancers treated with IV VAE (72). Patients were treated for an average of 48 days; with a maximum dose of 2000 mg, and mean survival was 130 days. Partial remission was seen in four patients, slowed

disease progression in two, progression of disease in two, and data was unavailable for two. Fever and fatigue were the most common side effects, with all side effects resolving after a treatment break. In the second study, a retrospective analysis was completed of matched-pairs for children with medulloblastoma treated with standard care, with or without anthroposophic medicine (including VAE). The study found no difference in 10year survival nor recurrence between the groups. The authors concluded that while treatment appeared to be safe, there was no survival benefit to be seen (73). Notably but not directly related to cancer but rather for safety considerations, mistletoe has also been used in children for other conditions, such as respiratory infections. For instance, in a study of 92 children with recurrent respiratory infections treated with VAE subcutaneous injections twice weekly for 5 weeks there was evidence of a positive immune response, reduced frequency of infections, and no safety concerns identified (74). While the evidence for benefit is thin in a pediatric cancer setting, available evidence indicates no safety concerns beyond what is known from adult populations. Given the potential for impact and low toxicity, selective use of mistletoe in a pediatric setting may be warranted.

# **Adverse Events and Side Effects**

VAE administered subcutaneously or intravenously is generally well tolerated (1, 2, 8, 23, 30, 44, 60, 75, 76). Overall, side effects are generally mild and self-limiting. Serious AEs have been documented but are rare. Certain side effects such as mild fever and local injection-site reactions may be considered desirable by some, as a surrogate marker for physiological response to treatment (23). Side effects of subcutaneous and IV applications differ and are discussed below.

# **Subcutaneous injections**

Side effects are common and expected, and mostly minor, dose-dependent, and self-limiting within a few days of treatment (2, 23, 68, 76). Common side effects include local reactions at the injection site (e.g.,

swelling, erythema, local pain, pruritus, induration, warmth), fatigue, mild flu-like symptoms, headache, mild fever, chills, flatulence and loose stools (2, 8, 23, 44). Localized reactions can sometimes appear at former injection sites for pre-exposed patients (2) and dose reductions might be required if reactions are severe (77). The side effect rate for mistletoe injections based on systematic reviews has ranged from 17.5% to 21.5%, with the vast majority being expected local reactions (77). More intense local skin reactions (>5 cm diameter) occur in less than 1% of cases (20) and are typically avoidable if a moderately progressive dosing approach is applied.

Reported serious adverse events are rare. They include urticaria and angioedema (37, 44), hypotension and loss of consciousness (78), anaphylaxis (<1%) (23, 78, 79), and severe delayed type hypersensitivity reaction (80).

<u>Common (>5%):</u> local injection-site reactions (e.g., swelling, erythema, pruritus, warmth, and induration).

<u>Rare (<5%):</u> fatigue, fever, chills, headache, flu-like symptoms, diarrhea/flatulence, and severe local reactions.

Rare but serious (1-4%): Angioedema, allergic reactions including anaphylaxis (<1%), hypotension and loss of consciousness, delayed hypersensitivity reaction, cellulitis at injection site.

#### **Intravenous infusions**

A phase I study investigated escalating doses (200-2000mg) in a variety of cancer types (60). The highest dose (2000 mg) was reported to have the same tolerability as the second lowest dose (400 mg). No serious AEs were deemed related to VAE. Adverse events related to VAE included allergic reactions, fever, weakness, eosinophilia and minor temporary ALT elevation. An observational study evaluated safety of IV VAE in 475 people (75). Twenty-two patients reported 32 adverse drug reactions, and none were serious. The most common was fever occurring in 8 people, followed by pruritus in 6. Other less common ADRs included

urticaria, inflammation of prior subcutaneous injection sites, vomiting, fatigue, infusion site irritation, myalgia, headache, paraesthesia, and rash. Compared to subcutaneous use, the ADR frequency of IV VAE was significantly lower (4.6% vs 8.4%, p = 0.005) mostly accounted for by the expected adverse skin reactions from SC injections. Iscador preparations had a higher frequency of ADRs compared with Helixor. Another retrospective observational study evaluated fever reactions in 59 patients receiving a total of 567 IV treatments (81). Forty-five (76%) of patients achieved a fever (>38.5°C) after at least 1 treatment, and fever was documented following 54% of infusions. Mean temperature increases following IV mistletoe was 1.5°C ± 0.8°C. Fevers were more common after Iscador infusions compared to other mistletoe products. Other adverse events were mostly fever-related (headache, shivering) in 48% of infusions, nausea in 15%, and allergic reaction in 0.6%. There were no grade 3 or 4 adverse events reported.

<u>Common (>5%):</u> Mild fever and related symptoms (headache, shivering), nausea

<u>Rare (<5%):</u> Pruritus, weakness, eosinophilia, minor temporary ALT elevation, urticaria, re-inflammation of prior subcutaneous injection sites, vomiting, fatigue, infusion site irritation, myalgia, headache, paraesthesia, rash

Rare but serious (1-4%): Allergic reaction (urticaria, angioedema).

# **Interactions with cancer treatments**

# Chemotherapy and radiotherapy

VAE has been studied alongside a variety of chemotherapy agents including carboplatin, gemcitabine, cyclophosphamide, 5-fluorouracil, methotrexate, and doxorubicin as outlined in Tables 1-4. None of these studies reported a worsening of treatment outcomes for survival, tumor response, or increased toxicity with the addition of VAE. As

discussed in the prior sections on efficacy, some studies reported better outcomes with the addition of VAE therapy. However, pharmacological studies to evaluate for interactions are lacking (23). A phase 1 pharmacokinetic study of VAE and gemcitabine found the combination was well tolerated, botanical/drug interactions were observed (52), but similar studies have not been performed for other chemotherapy agents. In vitro research corroborates the findings from human studies that have used VAE alongside chemotherapy without any worsening of treatment outcomes or toxicity. A study in 2017 found no induction or major inhibition of nine major cytochrome P450 isoenzymes with Helixor VAE products, making a clinically relevant pharmacokinetic herb-drug interaction unlikely (82).

Although direct pharmacokinetic and pharmacodynamic studies evaluating for interactions are lacking, the totality of evidence supports the premise that it is unlikely that there is any negative interaction with combined use with cytotoxic chemotherapy.

There is no known interaction of VAE with radiation therapy. Some studies in table 1 and 2 included people receiving radiation therapy without any negative interactions noted.

# Immunotherapy and targeted therapies

Due to the immunomodulatory properties of VAE, there has been some concern about the safety of combined use of VAE and immunotherapies and targeted therapies due to a theoretical additive effect. However, available evidence thus far has not demonstrated an increase in toxicity with combined use (83-87).

A multicentre observational trial evaluated the safety of targeted therapies with add-on VAE therapy compared to targeted therapy alone in 310 people (85). Targeted therapies included a variety of monoclonal antibodies (mAbs), immune checkpoint inhibitors (ICIs), and tyrosine kinase inhibitors (TKIs), but the majority of participants were using bevacizumab, rituximab, trastuzumab, or erlotinib. There was a significantly

lower AE rate in the combined group compared to control (20.1% vs 30.2%, p = 0.04) and a lower rate of discontinuation of standard oncology treatment in the combined vs control group (35% vs 60.5%, p = 0.03). A pilot study evaluated sixteen patients treated with ICI (Nivolumab, ipilimumab, pemprolizumab), of whom nine were treated with concomitant VAE (83). There was no statistically significant difference between groups with respect to AEs (67% in ICI plus VA, vs 71% ICI monotherapy). A retrospective study of 56 patients was conducted to evaluate the safety of combined mAb and intravenous Helixor VAE (84). Forty-three patients received combined therapy (defined as mAB and VAE administered on the same day), 12 received VAE therapy alone (no mAb within 1 month of VAE administration), and 8 received mAB therapy alone (no VAE within 1 month of mAb administration) (7 patients were included in more than one treatment group). The incidence of AEs was highest in the mAB monotherapy group (63%), followed by combined group (56%), and lowest in the VAE monotherapy group (42%). A multivariate analysis found increased odds experiencing an AE following mAB therapy compared to combined therapy (OR = 4.97, P = 0.008). Rates of serious AEs were similar for combined therapy (2%), mAB therapy (3%), and lower for VAE therapy (0.8%). Given the small number of people treated only with VAE or mAb, caution in interpretation is warranted. A small study of 15 patients with metastatic lung cancer treated with nivolumab alone (n=7) compared to nivolumab with VAE therapy (n=8) evaluated toxicity rates between groups (86). The toxicity rate in the nivolumab-alone group was 71.4% (5/7 participants) compared to 37.5% (3/8) in the combined group. An interim analysis of an ongoing prospective cohort study in patients with NSCLC evaluated the use of ICIs and VAE on symptom burden, QOL and OS and was published as a conference abstract. In an interim sample size of 20 within this study, the authors reported no clinically relevant increase in AEs due to VAE (87). Finally, a case report of a patient with metastatic stage IV clear cell renal cell carcinoma in the lung demonstrated no adverse effects from the combination of chemoimmunotherapy (interferon- $\alpha$ 2a, interleukin-2, fluorouracil, isotretinoin) and mistletoe treatment administered both IV and SC (66).

#### Other treatments

VAE injections were combined with radiofrequency ablation (RFA) in a case report with encouraging results (88). As noted below, when immunosuppressive treatments are applied, mistletoe use should be avoided.

## **Cautions and Contraindications**

Mistletoe should not be used by anyone with a known allergy or hypersensitivity to mistletoe. There is insufficient evidence regarding safety of mistletoe during pregnancy and lactation. Mistletoe should be used cautiously in people with autoimmune (AI) conditions although this is not a contraindication. Use should be avoided if immune suppressant medication is required to manage the AI condition due to the immunestimulating properties of mistletoe (2, 9, 13, 89). Given the need for immune suppression, mistletoe should not be used following a recent organ or bone marrow transplant. Mistletoe should be used cautiously in patients with brain tumors or metastases if there is unmanaged cerebral edema due to possible peri-tumoral inflammation with VAE, although evidence of harm from clinical studies is lacking (27). There is no clinical data or case reports using mistletoe for management of acute leukemias, however some suggest it should be considered a contraindication until more is known, given the possibility of leukocyte stimulation (23, 28). Although data from peer-reviewed sources is absent, there is some concern among practitioners about the use of fermented mistletoe products intravenously. The concern is that fermented products may increase the risk of allergic reactions, thus many clinicians use fresh unfermented aqueous extracts for IV use. There is an ongoing phase I clinical trial of IV fermented Iscador which should help to clarify whether there is any reason for concern (90).

#### **Autoimmune conditions**

Given the immunomodulatory properties of mistletoe, it has been theorized that it may exacerbate AI conditions. However, an uncontrolled observational study evaluated the safety of VAE therapy (IV, SC, IT) in people with

cancer with pre-existing AI conditions and failed to find an increased risk (91). In the cohort of 106 patients treated with VAE extracts, 17 patients (16%) experienced a VAE-related AE which is consistent with expected AE rate of other VAE-treated cancer patients. In a subgroup of 30 patients receiving long-term VAE therapy (>6 months), no exacerbations or flares of underlying AI disease were recorded. The most common AI conditions were Hashimoto's thyroiditis, psoriasis, ulcerative colitis, Grave's disease, and Sjogren's syndrome. Clinicians are recommended to discuss the theoretical possibility of AI condition flares with mistletoe use and consider the severity of the AI condition. It is recommended to not use mistletoe if the patient is using systemic immune suppressants to manage their condition.

#### **Brain tumors or metastases**

Many experts and VAE manufacturers recommend only using VAE in the absence of uncontrolled cerebral edema (27). The reason is due to the possible risk of peri-tumoral inflammation caused by mistletoe injections or infusions (27). There is no published data to confirm or refute this recommendation.

#### **Acute leukemias**

There is no published literature to demonstrate or refute a safety concern for VAE use in people with acute leukemia, however, some experts recommend caution based on the possibility of VAE stimulating the immune system (23, 27).

# **Dosing, frequency and length of treatment**

The maximum tolerated dose of IV VAE has not been established. In a phase I study, Helixor P (pine) was well tolerated up to the predefined maximum dose of 2000mg, with one dose limiting event occurring at this dose (60). IV mistletoe has been administered from 1-3 times weekly, over a duration of a few weeks to over a year in some observational studies. The optimal dose and length of administration is unknown.

The dose of subcutaneous injections varies based on VAE formulation, cancer stage, cancer type, and patient tolerance. It is typically recommended to use a dose escalation protocol starting with 0.01-1mg injections depending on the product, and increase based on tolerance. In Canada, Helixor (Viscosan) is the most common product; doses range from 0.1mg - 400mg, with administration most often 3 times weekly, and duration of use is most often several months (15, 17, 18, 37, 52). Although most clinical trials of VAE are a few months in duration, mistletoe has been used up to several years in observational studies and case reports without any apparent safety concerns (7, 46, 66, 70, 75, 76, 88, 92-94). In addition, long term usage of combined IV and SC VAE has been reported in case reports (67, 68).

Table 1: Clinical trials of subcutaneous (SC) mistletoe for cancer

Design   Measures	
al (2004) Ca Type: Q iii) Stable disease in 21 (84%) of participants which	
	lasted a median of
(50) Wictastatic Dosc, target 15 mg Survival 2.5 months.	lasted a median of
Colorectal Route: SC iv) Median survival 5.5 months.	
Cancer Admin: Toxicity v) Symptomatic relief observed in 10 (40%) of parti	cinants for nausea
Prior Tx: dose escalating, 3 (CTCAE) vomiting, diarrhea, constipation, fatigue and dyspne	
Chemotherapy injections a week until	
(resistant to toxicity or patient mild transient temperature elevation.	participants nau
5FU/LCV) bedridden	
Comparison:	
None None	
Piao et al Randomized N: 233 Agent: Helixor A Conventional chemotherapy QOL i) KPI scores significantly improved in the intervent	ion group
(2004) (37) Controlled Ca Type: Dose: (mixed type) (FLIC, KPI) compared to control (p=0.002).	ion group
Open label Breast, ovarian, 1-200 mg ii) Functional Living Index-Cancer (FLIC) scores significantly for the compared to control (p=0.002).	anificantly
NSCLC Route: SC Safety improved in the intervention group compared to con	
Stage: All Amin: Safety improved in the intervention group compared to control g	_
3 times weekly with dose the intervention group compared to 90 in control).	Toup (32 events in
escalation during  iv) One serious AE was noted in the VAE group: an	gioadama and
chemotherapy urticaria.	gioedema and
Comparison:	
control group receiving 4	
mg Lentinan injection	
daily	
Semiglasov Randomized N: 272 Agent: Lektinol 4 cycles CMF QOL i) 15 ng/0.5 ml given twice a week (30 ng/ml total) v	was found to be the
et al (2004) Placebo Ca Type: Breast, PS76A2 chemotherapy (cyclophosphamide, (EORTC QLQ-C30) dose which significantly improved QOL.	
(40) Controlled stage II/III Dose: methotrexate, fluorouracil) ii) Significant increase in CD4 count and CD4/CD8	ratio was observed
Double- Prior Tx: 10 or 30 or 70 ng/ml Adverse Events (p=<0.05).	
Blind Mastectomy Route: SC iii) VAE was very well tolerated, with local reaction	being the only
Admin: Immune markers adverse event related to the intervention.	
2x/week for 15 weeks	
during chemotherapy	
Comparison:	
placebo injection	
Semiglazov Randomized N: 352 Agent: Lektinol 4-6 cycles of CMF chemotherapy QOL (FACT-G, GLQ-1) FACT-G total score increased by 4.40±11.28 in M	E group, and
et al Placebo Ca Type: Breast, (PS76A2, an aqueous (cyclophosphamide, methotrexate, 8, Spitzer's uniscale) decreased by 5.11±11.77 in placebo (p<0.0001).	
(2006)(42) Controlled stage II/III mistletoe extract) fluorouracil) ii) GLQ-5 sub-score was significantly better (lower)	in ME compared
Safety (Adverse events) to control group (42.9±125.0 vs 60.3±94.0 p<0.0001	•

	Double- Blind		Dose: 15 ng mistletoe lectin/0.5 ml Route: SC Admin: 2x/week for 4-6 cycles of chemotherapy Comparison: placebo injection			worsened in both groups but moreso in placebo group than ME group (p = 0.0007).  iii) Spitzer's uniscale improved in ME group compared to placebo (12.2±30.7 vs 10.8±26.1 p<0.0001).  iv) Well tolerated, local reactions occurred in 17.6% of participants.
Enesel at al (2005) (21)	Randomized Controlled	Ca Type: mixed gastroesophageal and abdominal cancers (esophageal, gastric,	Dose: 60 mg/ml Route: SC Admin: every second day from 2 weeks before to 2 weeks after surgery Comparison:	Surgery	Cellular Immunity (CD2, CD3, CD19, CD4, CD8, NK)  Humoral Immunity (IgG, IgA, IgM, complement)  QOL (KPS)	i) Compared to controls, treatment arm had significantly higher: WBC counts before and after surgery (p < 0.001), lymphocytes after surgery (p < 0.001), complement post-surgery (C3 and C4) (p < 0.001), immunoglobulins post-surgery (particularly IgA and IgM), (p<0.05), CD4/CD8 ratio before and after surgery (p<0.05), and NK cell levels significantly increased overall (p<0.001).  ii) KPS score significantly increased in the intervention group (p<0.01) compared to a significant decrease in the control group (p<0.05).
Troger et al (2009) (20)	Randomized Controlled Open	metastatic breast	<b>Dose:</b> 0.01-5 mg	6 cycles CAF chemo	QOL (EORTC QLQ-C30) Neutropenia	i) Mean differences were significantly better for 12 of the 15 QOL endpoints in the mistletoe group compared to control (range: p= 0.017 to p<0.001). Clinically relevant changes (5-point differences) were noted for 9 QOL endpoints.  ii) Neutropenia occurred non-significantly less in the intervention group compared to control (p=0.182).
Reif M et al (2019) (26)	See Troger	et al (2009) (19) as	s above (re-analysis of data the abstract was available	a for additional outcomes). *Only e		i) Absolute T4, monocyte, and absolute NK cell counts, and absolute T8 cell counts were correlated with CRF with statistical significance (p $\leq$ 0.05) or tendency (0.05 < p < 0.1). in the control arm. However, these correlations in the Iscador M arm were weaker and not significant. May indicate that VAE attenuates inflammatory immune response which contributes to effect on CRF.
al (2010)	Randomized Controlled Open	Ca Type: Stage I/II breast, post- treatment	8	None, VAE was initiated SC 2 weeks post-treatment completion	Cytokines (IL2, IL4, IL6, IL10, TGF-b, IFN-y)	i) Concentrations of IL6 and IFN-y significantly increased from baseline after treatment compared to control (p=0.013 and p=0.009, respectively).  ii) No significant changes from baseline were noted for IL2, IL4, IL10, TGF-b.

			aamplating aanaar		I	
			completing cancer			
			treatment (surgery,			
			chemo radiation)			
			Comparison:			
			standard treatment			
			alone			
Kim et al	Randomized		Agent: abnobaVISCUM	5-DFUR (chemo)	-	i) QOL: Compared to control, the following improved in the mistletoe
(2012)(19)	Controlled	Са Туре:	"Q"		(EORTC QLQ-C30,	group: global health status (p=0.0098), pain (p=0.038), eating
	Open Pilot	Gastric (stage Ib	Dose:		ST022)	restriction (p=0.037), and hair loss (p=0.023).
		primarily)	0.02 mg- 20 mg			ii) Significantly higher WBCs (p=0.0101) and eosinophil counts
		Prior Tx:	Route: SC		Liver Function	(p=0.0036) were observed in the intervention group.
		Surgery	Admin:			iii) No differences were noted for CD16/CD56, CD19 lymphocytes,
			dose escalating, 3X/week		Immune Markers	TNF-a and IL2.
			beginning 7 days after		(TNF-a, II2,	iv) No serious AEs attributed to mistletoe.
			surgery, for 24 weeks.		CD16/CD56, CD19	
			Comparison:			
			standard treatment			
			alone			
Bar-Sela,	Phase II,	N: 72	Agent: Iscador Q	Carboplatin-based combination	Toxicity	i) Control group had more chemotherapy dose reductions (44% vs 13%
2013 (32)	randomized		-	chemotherapy given in 21-day	(CTCAE)	p = 0.005).
, ,		NSCLC	Route: SC	cycles	,	ii) Treatment group had fewer grade 3-4 non-hematological toxicities
			Admin:		Quality of life (EORTC	(41%  vs  16%, p = 0.043),  hospitalizations  (54%  vs  24%, p = 0.016),
		` .	dose escalation from 0.01		QLQ-C30 and QLQ-	and rate of peripheral neuropathy (p=0.03).
		adenocarcinoma)	to 10 mg of mistletoe,			iv) No difference in grade 3-4 hematological toxicity or total grade 3-4
			given every other day		,	toxicity (48% vs 57%, NS).
		IIIA-IV (majority			Tumor response	v) No difference in primary QOL questionnaires.
			chemotherapy alone		(RECIST criteria)	vii) mOS in both groups was 11 months.
		Prior Tx:	1,5		,	viii) Median TTP was 4.8 months for control vs. 6 months in iscador
		No prior chemo			Overall Survival	(NS).
Mansky,	Phase I		Agent: Helixor A	Stage I:	CT scan -baseline and	i) 112 AEs attributed to mistletoe. Most common: injection site
	Uncontrolled	Ca Type: Mixed		Gemcitabine dose (750 mg/m²) IV		reaction (42 events), localized induration (20 events), grade 1-2 non-
, ,	2 Stage	V -	Stage I: Escalating dose	on day 1 & 8 of a 3-week cycle		neutropenic fever (22 events) and grade 1-2 flu-like symptoms (10
	Design		1mg – 250mg		Adverse Events	events). 2 grade 3 events - cellulitis at injection site
	-		Stage II: Dose right	Stage II:		ii) MTD was 250 mg for mistletoe.
				Escalating IV gemcitabine (20%	(	iii) Mistletoe did not affect gemcitabine pharmacokinetics. Clinical
		_	Route: SC	increments) dosing	Lab Values	response similar to gemcitabine alone.
			Admin:			iv) 33 completed 3 cycles. 6% achieved partial response, 42% achieved
			Stage I: Dose escalation		Clin. Eval.	stable disease and 43% progressed (9% not evaluable).
		-	of mistletoe, fixed dose			v) All developed ML-3 IgG antibodies, with higher levels achieved
			gemcitabine		MTD & DLT	with increasing doses of mistletoe. Cytokines were not affected.
			Stage II: Fixed dose			<u> </u>
			mistletoe, escalating		Survival	
			gemcitabine			
					Clinical Response	

Troger,	Phase III	N: 220	Agent: Iscador Q	Standard supportive care only	Overall Survival	i) mOS was 4.8 months in the intervention group compared to 2.7
2013 (33)	Randomized	Са Туре:	Dose: escalating dose		QOL	months in control group (HR: 0.49, 95% CI: 0.36-0.65, p<0.0001).
	Controlled		(0.01 mg - 10 mg)	No anti-neoplastic therapies	Vital Signs	ii) No adverse events related to mistletoe, and fewer AEs in treatment
	Open-Label		Route: SC	provided	Performance Status	vs control group (17 vs 53 respectively)
	1	Stage:	Admin:		Weight	iii) Frequency and severity of symptoms were significantly lower in the
		_	3X/week up to 12			intervention group compared to control for pain (p<0.0001), weight
		` ′	months		Medication Use	loss (p<0.0001), energy (p<0.0001), nausea/vomiting (p<0.0001),
			Comparison:		Safety	diarrhea (p=0.0033) and anxiety (p=0.046).
			supportive care only		(CTCAE)	1 , , , ,
		2-4 (n=108)	,			
		Prior Tx:				
		205 had surgery				
-	-		n 96 patients in the mistle	toe group and 72 patients in the	QOL and symptoms	Compared to control, Iscador Q:
	control group	).			(EORTC QLQ-C30)	i)Had improved global health and functional scales.
(38)						ii) Improved symptom scale in 6 out of 9, including pain (95% CI: -29
					Body weight	to –17), fatigue (95% CI: –36.1 to –25.0), appetite loss (95% CI: –51 to
						−36.7), and insomnia (95% CI: −45.8 to −28.6).
						iii) increased body weight (5.3% increase vs 3.2% decrease, p<0.001).
Reif et al	See Troger, 2	013 (33) as above	(post-hoc analysis)		Pain (EORTC QLQ-	i)Patients in the control group received more potent and frequent
,2019 (39)					C30) and consumption	analgesics than those in the VAE group (OR = 0.005, 95%-CI [0.001;
					of analgesics	0.014]).
						ii) Post-baseline pain EORTC QLQ-C30 scores were lower in the
						VAE arm than in the control arm: mean OR = 0.013, 95%-CI [0.006;
						0.028]).
						iii) investigators reported lower pain levels in VAE group (mean OR =
						0.034, 95%-CI [0.009; 0.123]) than in the control group.
_	Randomized	N: 20	Agent: Iscador P	None	1-year PRDFS	i) 1-year PRDFS was 55.6% in mistletoe arm compared to 12% in
2014 (34)	Controlled	Са Туре	Dose:		(primary)	historical controls (p=0.0041, 95% CI: 21.2%-86.3%). The rate in the
	Open-Label	Relapsed	escalating dose (0.01 mg			etoposide group was 27.3% compared to 12% in historical controls
		Osteosarcoma	- 20 mg).		Quality of Life	(p=0.2724, 95% CI: 6.0%-61.0%).
		Stage:	Route: SC		(EORTC QOL-C30,	ii) The median PRDFS at the time of analysis was 39 months in the
		1 stage 1B	Admin:		PedsQL)	mistletoe group (range 2-73 months) and 4 months in the etoposide
		14 stage IIA/B	3X/week for 12 months			group (range 1-47 months), no statistical analysis applied, however the
		_	Comparison:		Safety	follow up was ongoing.
			oral etoposide daily for		(CTCAE)	iii) Compared to baseline, mistletoe therapy significantly improved
		Prior surgery and	21d of 28d cycle (total of			QOL measures of physical functioning (p=0.046), emotional
		chemo, no prior	6 cycles)			functioning (p=0.014), social functioning (p=0.003), global health
		radiotherapy.	(historical controls were			(p=0.013), fatigue (p=0.005), pain (p=0.012), dyspnea (p<0.0001),
			also used to evaluate			insomnia (p=0.020) and financial strain (p<0.0001).
			each treatment arm)			iv) No toxicity was noted for VAE other than minor local erythema
						after injection and hypotension in one patient.
1						

Longhi,		S	See Longhi, 2014 (34), as a	bove	PRDFS (long-term	i)The mistletoe arm saw a median PRDFS of 106 months compared to
2020 (53)					follow up)	7 months in the etoposide arm (HR 0.287, 95% CI 0.076-0.884, p =
					-	0.03). 5 of 9 patients never relapsed in the VAE arm, compared to the
						etoposide group in which all patients relapsed.
						ii)Through a model, the estimated 10-year overall survival rates were
						64% and 33% in the mistletoe and etoposide arms, respectively
						(statistical significance not calculated).
Troger,	Randomized	N: 65	Agent: Helixor A	Adjuvant chemotherapy	Quality of Life	i) Compared to control, mistletoe improved QOL from baseline
2014 (17)	Open-Label	Ca Type: Non-	Dose:	(6 cycles CAF)	(EORTC QLQ-C30)	significantly more for role function (p<0.001) emotional function
		metastatic Breast	escalating dose of 1 mg-			(p<0.001), social function (p<0.05), cognitive function (p<0.01), pain
		Prior Tx:	50mg		Neutropenia	(p<0.001), anorexia (p<0.001), diarrhea (p<0.001), insomnia (p<0.05),
		Surgery	Route: SC		(neutrophil count)	nausea/vomiting (p<0.001), and constipation (p<0.05).
			Admin:			ii) Compared to control, mistletoe did not improve QOL parameters
			3X/week during 6 cycles		AEs	from baseline for global health, physical function, fatigue, dyspnea and
			of chemotherapy		(CTCAE-v3)	financial strain .
			Comparison:			iii) No significant change in neutropenia occurrence (p=0.628).
			chemotherapy alone			iv) Overall VAE was well tolerated. The only notable adverse events
						were erythema >5 cm (42 events, 2.7% of injections), and. one
						participant experienced rhinoconjunctivitis and withdrew from the
						study.
Pelzer,	Randomized		Agent: Helixor A or	CAF chemotherapy	Temperature	i) 2 fevers observed, neither were long-lasting.
2018 (18)	Controlled	~ I	Iscador M	(6 cycles)		ii) No significant differences in neutropenia between groups (p=0.178)
	Open-Label		Dose: Helixor A,		Neutropenia	iii) Compared to control, mistletoe significantly improved role
		Breast	escalating dose of 1 mg-			functioning (p<0.0001), emotional functioning (p=0.0226), pain
			50 mg		Quality of Life	(p<0.0001) and diarrhea (p=0.0311).
		Surgery	OR		(EORTC QLQ-C30)	iv) Compared to control, mistletoe did not significantly affect global
			Iscador M: escalating			health status, physical functioning, cognitive functioning, social
			dose of 0.01 mg, 0.1 mg-		Relapse	functioning, fatigue, nausea/vomiting, dyspnea, insomnia, appetite loss,
			5 mg		(5 year follow-up)	constipation and financial difficulties.
			Route: SC			v) Other than local skin reactions, no AEs were observed for mistletoe
			Admin:		Metastasis	therapy.
			3X/ week during 6 cycles		(5 year follow-up)	vi) 56/65 tx group and 29/31 controls were evaluable for DFS. 15/56 in
			of chemotherapy.			tx arm developed relapse or metastasis compared to 8/29 controls
			Stopped within 3 weeks			(p=0.76). Median DFS could not be calculated.
			of chemo			
			discontinuation			
			Comparison:			
			chemotherapy alone			
Add: addition	onal Admin.	administration Al	F. adverse event Co. con	er CAE evelophosphamida/dox	orubicin (Adriamycin)/flu	orouracil Chemo: chemotherapy Clin. Eval: clinical evaluation CMF:

Add; additional, Admin; administration, AE; adverse event, Ca; cancer, CAF; cyclophosphamide/doxorubicin (Adriamycin)/fluorouracil, Chemo; chemotherapy, Clin. Eval; clinical evaluation, CMF; cyclophosphamide/methotrexate/fluorouracil, CRF; cancer related fatigue; CTCAE; common terminology for adverse events, CT; computerized tomography, DFUR; Docetaxel/epirubicin/doxifluridine, DLT; dose limiting toxicities, EORTC QLQ-C30; European Organization for the Research and Treatment of Cancer Quality of Life Questionnaire, KPI; key performance indicators, KPS; Karnofsky performance status, LCV; leucovorin, ML; mistletoe lectin, MTD: maximum tolerated dose, N; number of participants NR; not reported, NS; non-significant, NSCLC; non-small cell lung cancer, PRDFS; Post-Relapse-Disease-Free-Survival, QOL; quality of life, Rad; radiation, SC; subcutaneous, Surg; surgery, Tx; treatment, VAE; Viscum album extract, yoa; years of age, 5-FU; fluorouracil

**Table 2: Clinical trials of intravenous mistletoe for cancer** 

Reference	Study design	Participants	Intervention	Concomitant	Outcomes and	Results
				treatment	measures	
Cazacu et al	Randomized	N: 64	Agent: Isorel	Chemotherapy	Survival	i) 4 treatment AEs in the surgery + chemotherapy
(2003) (61)	Controlled	Ca Type:	Dose:	(5-FU)		group compared to none in the surgery +
	Open	Advanced	5 mg/kg in saline infusion (500 ml)			chemotherapy + mistletoe group.
		colorectal	Route: intravenous			ii) Median survival was significantly better in the
		Prior Tx:	Admin: 3 infusions weekly after surgery			mistletoe group compared to the surgery +
		Surgery	alongside adjuvant chemotherapy			chemotherapy alone group (p< 0.05).
			Comparison groups:			
			Surgery alone (no adjuvant treatment),			
			surgery + adjuvant chemotherapy			
Huber et al,	Phase I	N: 21	Agent: Helixor P	None	MTD	i) 0 drop outs. One DLT occurred at the 2000 mg
2017 (60)	Safety Study	Ca Type: mixed	Dose:			dose - generalized urticaria allergic reaction
		Stage: advanced/	Phase I dose finding design: 200mg, 400 mg,		DLT	requiring IV anti-histamines.
		metastatic	700 mg, 1200 mg and 2000 mg		(AE >/= grade	ii) Tolerability of 2000 mg did not differ from 400
		Prior Tx:	Route: Intravenous		2)	mg.
		15 Surgery	Admin: 1 infusion weekly for 3 weeks. A			iii) 6 serious AEs occurred during the study, none
		14 Chemotherapy	3+3 dose design was implemented until the		Safety	attributed to mistletoe.
		9 Radiotherapy	maximum dose (2000 mg). If the max dose		(CTCAE,	iv) 25 AEs were deemed possibly related to the
		4 Immunotherapy	was achieved, it was applied for 9 more		physical exam,	intervention (all occurring at 2000 mg dose).
			weeks		blood work)	Allergic reaction (1), grade 1 fever (4), weakness
			Comparison:			(3), eosinophilia (2), and temporary minor ALT
			Phase 1 internal comparison - Safety of		Tolerability	elevation (2).
			different mistletoe infusion doses			v) 2 patients had unexpected temporary tumor
						marker improvement. One patient had a slowed
						progression.

AE; adverse event , Admin; administration, Adv/mets; advanced and/or metastatic disease, ALT; Alanine-transaminase, Ca; cancer, CTCAE; common terminology for adverse events, DLT; dose limiting toxicity, MTD; maximum tolerated dose, , temp; temperature, Tx; treatment, WBC; white blood cell count, 5-FU; fluorouracil

<u>Table 3: Clinical trials or observational studies of intratumoral, intravesicular, intrapleural, or transcatheter mistletoe</u>

Reference	Study design	Participants	Intervention	Concomitant treatment	Outcomes and measures	Results
Elsasser-	Phase I/II	N: 30	Agent:	None	Recurrence	i) No local or systemic side effects noted.
Beile et al		Ca Type: Bladder	aqueous mistletoe extract		(Cytology,	ii) At the 12-month mark, 30% developed recurrence. No clear
(2005) (95)		Prior Tx:	Dose:		ureterocystoscopy)	association between dosage and recurrence rate was found.
		Transurethral	10-5000 ng/ml			iii) Recurrence rate was comparable to historical controls.
		resection	Route:			
			intravesicular			
			Administration:			
			6 weekly instillations. Extract			
			retained 2 hours in bladder.			
Bar-sela et al	Open	N: 25 (23	Agent: Iscador M	Peritoneal	Drainage Time	i) Paracentesis interval was 7 days prior to mistletoe, and extended to
(2006) (96)		evaluable)	Dose: 10 mg diluted in 10-15 ml	puncture	Intervals	12 days after the first instillation (p=0.001).
		Ca Type: mixed	saline			ii) No differences in abdominal circumference, volume drained or
		stage IV cancers,	Route:		Abdominal	symptom scores noted. Transient abdominal pain was noted in one
		mostly	peritoneal catheter used for		Circumference	participant for 1 hour which self-resolved. No other AEs were noted
		gastrointestinal	drainage (injection)			during the trial.
			Admin: following abdominal		Drainage Volume	
			punctures for drainage			
			Comparison:		Symptoms	
			previous drainage parameters			
Gaafar, 2014	Randomized	N: 23	Agent: Viscum Fraxini-2	Fluid drainage	Physical Exam	i) Overall clinical response was 61.5% in the mistletoe group and 30%
(97) Gaafar,	Controlled	Ca Type: lung	<b>Dose:</b> 5 ampoules in 10 cc			in the bleomycin group, however the difference was not significant
2014 (97)		(mixed types)	glucose 5%		Chest Radiography	(p=0.21).
			Route: intrapleural, via chest		(Pleural effusion	ii) Adverse events reported in the mistletoe group included fever, chills,
			tube		evaluation)	headache, malaise and allergic reaction (requiring discontinuation and
			Administration:			steroid injection). No hospitalization was required for any of the
			up to once weekly for 6 weeks if		Adverse Event	adverse events.
			needed until dryness of pleura		(CTCAE v4.0)	
			Comparison:			
			bleomycin (60 units) once			
			intrapleurally			
Rose et al,	Phase Ib/IIa	<b>N:</b> 36	<b>Agent:</b> Abnoba viscum Fraxini 2	None	Safety	i) No dose limiting toxicity was found up to 675mg.
2015 (98)		Ca Type: Bladder	<b>Dose:</b> range from $45 - 675$ mg			ii) A total of 214 AEs were reported, 76 were deemed possibly or
		Cancer	Route: intravesicular		Recurrence	probably related to intervention. Most common were local skin
		Prior Tx: Surgery	Administration: weekly for 6			reaction, urinary tract infection, and pyrexia. All participants recovered
		(transurethral	weeks, dose escalating to find			fully.
		resection)	tolerable dose.			iv) Based on 30 evaluable patients, at the 12 week mark, 66.7% had no
						visible "marker" tumor (remnant of tumor purposely left over after

						surgery to assess intervention) remaining and negative biopsy. Based on 19 evaluable participants, the recurrence rate was 26.3%.
Cho et al, 2016 (11)	Open-Label Phase III Single Arm Multicenter	N: 62 Ca Type: mixed. Large proportion were lung cancer	Agent: Abnovaviscum Dose: 20 mg Route: direct injection into pleural space Administration: after pleural effusion drainage, injection administered with dosing schedule based on newly- generated pleural effusion	Pleural effusion drainage	Pleural Effusion  QOL (KPS score)  Safety	<ul> <li>i) Complete pleural effusion response rate 79.0%, compared to historical reference of 64.0% (p &lt;0.0001).</li> <li>ii) No significant changes in KPS scores were noted compared to baseline.</li> <li>iii) 309 AEs occurred. 42 could not be excluded as causal with intervention; most frequent were localized reaction, pyrexia, chills, fatigue and pain. All AEs fully resolved. 2 serious AEs occurred that could not be excluded which included serious pleuritic and pain in one patient.</li> </ul>
Galun et al, 2019 (99)	Conference abstract: Prospective cohort analysis	N: 107 Ca Type: non-resectable hepatocellular carcinoma	Agent: Iscador Qu Dose: unknown Route: hepatocellular transcatheter Administration: unknown	Lipitol and cisplatin	Survival time	i) A significantly better median survival time was found in the mistletoe group who received Iscador Qu in addition to standard treatment, compared to the control group, at 430 and 246 days, respectively (HR = 0.36; CI 95%: 0.23-0.57).  ii) Participants in the mistletoe group who developed a fever had a slightly better survival time than those who did not, though the difference was not statistically significant.
Lee et al (2019) (100)	Retrospective	N: 52 Ca type: Lung Cancer Stage: With malignant pleural effusion	Agent: Helixor M Route: Pleural Catheter (pleurodesis) Dose: 100mg, if ineffective the dose increased by 100mg each instillation Administration: 1-5 treatments as needed (every other day for repeat instillations) Comparison: None	Drainage catheter	Malignant pleural effusion control Safety	<ul> <li>i) The one month recurrence rate of malignant pleural effusion was 48%.</li> <li>ii) 25% of patients experienced pain related to the procedure and 15% had fever &gt;38 °C.</li> </ul>

Ca; cancer, Tx; treatment, AE; adverse event, CTCAE; common terminology for adverse events, KPS; Karnofsky performance status, NS; non-significant, QOL; quality of life

Table 4: Observational research of subcutaneous or IV mistletoe for cancer

Reference	Study design	Participants	Intervention	Concomitant Tx	Endpoints and	Results
Pussing at al (2007)	Dungmantive	N: 71	Agent: Iscador	None	Measures Immune Effects	Swift escalation of dose resulted in more local reactions
Bussing et al (2007) (77)	Prospective Cohort	Ca Type: Breast, Prostate, Colorectal Stages: I-IV	Agent: iscador Dose: 0.01mg – 20mg Route: SC Administration: 2x/week, over 6 months Comparison: slow incremental increase vs. rapid dose escalation	None	QOL	<ul> <li>Swift escalation of dose resulted in more local reactions compared to slow incremental increase.</li> <li>No differences were noted between groups regarding body temperature and QOL.</li> <li>No differences between dosing schedules were noted for CD3, CD4, CD8 or CD4/CD8 ratio.</li> <li>Swift escalation group had a significant decrease in HLA-DR+ T-Cells compared to a slight increase in the slow escalation group (p &lt; 0.05).</li> </ul>
Beuth et al (2008) (46)	Retrospective Cohort	N: 681 (167 mistletoe, 514 control) Ca Type: Breast Stages: I-III	Agent: Helixor Dose: not specified Route: not specified Administration: frequency not specified, used for up to 5 years post-cancer treatment Comparison: No mistletoe	Standard cancer treatments (surgery +/- chemotherapy, radiation, endocrine therapy)	Safety during aftercare (post-cancer treatment) (medical records)  Symptoms (obtained from medical records) during aftercare (post-cancer treatment)	<ul> <li>Adverse drug reactions to mistletoe in the treatment group were 10% (local reactions, erythema, pruritus, flu-like symptoms, one case of generalized reaction).</li> <li>In the aftercare period (after surgery, chemo, radiation were completed), disease or treatment-related symptoms were significantly lower in the mistletoe vs control group ( 56.3% vs 70%, P &lt; 0.001).</li> <li>Adjusted odds ratio of symptoms for mistletoe treated group was 0.51 (95% CI: 0.32-0.81).</li> <li>There was no difference between groups for rates of relapse, metastases, or death.</li> </ul>
Bock et al (2014) (50)	Retrospective	N: 324 Ca Type: Colorectal Stage: non- metastasized CRC, stages I-III	Agent: Iscador Q Dose: total 16 to 20mg per week Route: SC Administration: daily doses were left up to physician's discretion Comparison: NA	Chemotherapy or radio-chemotherapy	Cancer Related Fatigue	- Those who received mistletoe in addition to standard care had a cancer-related fatigue rate of $8.8\%$ compared to $60.1\%$ in the control group (p < $0.001$ ).
Schad et al (2014) (101)	Retrospective	N: 39 Ca Type: Advanced Inoperable Pancreatic Cancer Stage: II-IV	Agent: Helixor, Abnoba Dose: escalating doses up to 160mg (Abnoba) or 1400mg (Helixor) Route: intratumoral Administration: alternately to chemotherapy in 4-week intervals or more Comparison: NA	Chemotherapy	Safety Survival	<ul> <li>No serious intervention-related adverse effects. Increased body temperature was seen in 14% and fever in 11%.</li> <li>Median survival 11 months (11.8 for stage III and 8.3 for stage IV).</li> <li>Considered feasible, well-tolerated and safe.</li> </ul>

Steele et al (2014) (76)	Observational	N: 1923 Ca Type: multiple types Stage: 0-IV	Agent: mixed  Dose: varied, ≤0.02 to 60mg  Route: SC  Administration: varied, most often 3X/week, median length of mistletoe therapy 4.6 months  Comparison: NA	Conventional care	Safety: AEs & ADRs	<ul> <li>21.5% experienced either an expected effect or an adverse drug reaction.</li> <li>264 ADRs in 162 patients (8.4%). 42.1% were possibly related, 53.4% were probably related and 4.5% were certainly related to mistletoe treatment.</li> <li>ADRs included: local skin reaction &gt;5cm, &gt;38 C temp, chills, fatigue and malaise. 50.8% of ADRs were classified as mild and 45.1% moderate.</li> <li>11 severe ADRs which included 8 patients with temp &gt;40C for less than 24 h, 1 with severe injection site swelling, 1 with general urticaria and 1 with syncope. All patients fully recovered.</li> <li>No life threatening ADRs occurred.</li> <li>ADRs in general appeared lower with the combination of mistletoe therapy and conventional care.</li> <li>Mistletoe ADR rate increased as dose increased.</li> </ul>
Steele et al (2014) (75)	Retrospective	N: 475 Ca Type: multiple types Stages: I-IV	Agent: Helixor, Abnoba, Iscador Dose: ranged 10 to 400mg Route: IV and SC Administration: mixed Comparison: NA	Conventional care	Safety: AE's & ADRs	<ul> <li>No serious ADRs occurred.</li> <li>22 patients reported 32 ADRs (59.4% mild, 40.6% moderate).</li> <li>Iscador brand showed relative higher frequency of ADRs compared to the other products.</li> <li>Intravenous mistletoe had significantly less ADRs than subcutaneous administration (4.6% vs 8.4%, p=0.005).</li> </ul>
Steele et al (2015) (102)	Retrospective	N: 123 Ca Type: multiple types Stage: mixed and some unknown, but 47.2% stage IV	Agent: Helixor, Abnoba, Iscucin Dose: 0.02 to 250mg, median dose 60mg Route: intratumoral Administration: varied, majority received 2-6 applications, up to 1 month Comparison: NA	Mixed (SC, IV, both)	Safety: AE's & ADRs	<ul> <li>26 patients experienced a total of 74 ADRs (21.1%).</li> <li>Most common ADRs were body temperature increase or immune related effect, of which 83.8% were mild and 14.9% moderate.</li> <li>One possible severe ADR occurred (hypertension) with no serious ADRs occurring.</li> <li>Intratumoral ADR rates were 3x higher than SC and 5x higher than intravenous application rates when compared with external data.</li> </ul>
Von Schoen-Angerer (2015) (93)	Retrospective Case-series	N: 8 Ca Type: Bladder Cancer Stage: Majority were non-muscle invasive cancer.	Agent: Iscucin Salicis Route: SC Dose: strengths F (0.125mg), G (2.5mg) and H (50mg) Administration: varied from 1x/week to daily based on fever and inflammatory reactions Comparison: NA	Mixed	Recurrence	<ul> <li>Median tumor-free duration was 48.5 months.</li> <li>High dose mistletoe showed possible benefit in 5 of 8 patients, 2 patients could not be assessed and 1 showed uncertain effects of mistletoe.</li> <li>No tumor progression was observed in any of the 8 patients.</li> <li>No patient stopped treatment due to intolerance/side-effects.</li> </ul>
Sunjic et al (2015) (92)	Retrospective Case-report series	N: 74 Ca Type: multiple Types Stage: majority were advanced stages	Agent: Isorel (A, M & P)  Dose: not reported, as per manufacturers guidelines  Route: SC, IM, IV  Administration: 3X/week first year after diagnosis, then	Conventional care (primarily surgery and radiation)	Clinical Effect (not adequately described)	<ul> <li>There was no tumor recurrence in 47% of cases, partial cancer regression in 38% of cases, and no cases of worsening condition.</li> <li>Not much can be stated from this study due to poor methodology.</li> </ul>

			maintained or reduced to 1X/week in cases of remission Comparison: NA			
Axtner et al (2016) (103)	Retrospective	N: 240 Ca Type: Advanced Pancreatic Cancer Stage: stage IV	Agent: mixed Dose: not reported Route: SC (89.2%), IV (35.2), intratumoral (19.3%) Administration: alongside chemotherapy, durations not reported Comparison: chemotherapy only and VA only	Chemotherapy	Feasibility Survival	Patients receiving >4 weeks of mistletoe in addition to chemotherapy had longer survival compared to those who only had chemotherapy (12.1 vs 7.3 months) (log rank test, X2=6,p=0.014).  Patients receiving VA only had longer survival than those receiving neither chemotherapy nor VA therapy (5.4 compared to 2.5 months) (log rank test X2 = 7.6, p=0.006).
Schad et al (2017) (104)	Retrospective	N: 1361 Ca type: Multiple types Stage: varied	Agent: Abnobaviscum Fraxini (44%), Mali (22.3%), Quercus (22.1%), other (11.6%)  Route: SC  Administration: duration not reported  Comparison: low initial dose group ≤ 0.02mg (516 patients) vs. high initial dose group >0.02mg (845 patients)	Not reported	Safety: AEs & ADRs (high vs low starting dose)	<ul> <li>Initiation of a high dose was associated with a significantly higher risk of ADR compared to initiation of treatment with low dose (20.7% vs 0.8%, p≤ 0.001).</li> <li>No serious ADRs occurred.</li> </ul>
Schlappi et al (2017) (81)	Retrospective	N: 59 Ca type: Multiple types Stage: 59% advanced or metastatic disease	Agent: most frequently used was Iscador M Dose; varied Route: IV Administration: varied considerably Comparison: NA	None	Fever (≥ 38.5 C°) Safety (CTCAE v 4.0)	<ul> <li>Out of 59 patients, receiving a total of 567 intravenous infusions, 45 patients (76%) achieved a fever after at least 1 treatment.</li> <li>Mean temperature increase 1.5 C<sup>0</sup> +/- 0.8 C°.</li> <li>No AE's over grade 2 occurred. One grade I allergic reaction occurred.</li> </ul>
Thronicke et al 2017 (83)	Retrospective	N: 16 Ca type: Primarily lung cancers (69%) Stage: IIIA/IV(Progressive or metastatic)	Agent: Varied: Abnobaviscum,Helixor P Iscador Q Dose: varied Route: Varied (SC or IV or both) Administration: median duration was 84 days (range of 1-196 days) Comparison: ICI alone	Immune checkpoint inhibitors (ICI)	Response Rate  AEs (CTCAE)	<ul> <li>AE frequency rate was 68%, with 11 participants experiencing at least 1 AE.</li> <li>No grade 3 or 4 AEs occurred.</li> <li>Most frequent AEs reported were malaise, pyrexia, bronchitis and skin reaction.</li> <li>Multivariate regression showed no significant association between the combination of mistletoe and immunotherapy for AE rate (OR: 1.467, 95% CI: 0.183-11.693, p=0.720).</li> <li>Progressive disease was observed in 71.7% of participants in the immunotherapy alone group, compared to 44.4% in the combined treatment group (p=0.36). Stable disease was observed in 28.6% of participants in the immunotherapy alone group, compared to 22.2% in the combined treatment group (p: not available). Overall, no statistically significant differences were found between groups.</li> </ul>
Fritz, et al (2018) (105)	Retrospective Case-	N: 18,528 Ca type: Breast	Agent: Lectinol <sup>R</sup> , Abnoba, Helixo, Iscador, and	Standard breast cancer treatment	Survival	- Multiple types of mistletoe preparations, doses, administrations, etc.
	Controlled	Cancer	Aviscumine		QOL	

		Stage: Moste were I or II	Dose: not reported Route: variable and uncertain Administration: not reported Comparison: Standard breast cancer treatment alone			<ul> <li>No survival benefit when mistletoe is added to conventional treatment.</li> <li>No QOL benefit observed when mistletoe compared to conventional treatment.</li> </ul>
Schad et al (2018) (58)	Retrospective	N: 158 Ca type: NSCLC Stage: IV	Agent: Abnobaviscum, Helixor and Iscador Dose: Not reported Route: SC, IV, intratumoral Administration: Not reported Comparison: Chemotherapy alone	Chemotherapy	Survival	<ul> <li>Median survival for patients receiving mistletoe + chemotherapy was 17.0 months compared to 8.0 months in the chemotherapy group alone (p=0.007).</li> <li>Overall survival was significantly prolonged in the mistletoe combination group (HR: 0.44, 95% CI: 0.26-0.74, p=0.002).</li> <li>1-year survival was 60.2% in mistletoe group compared to 35.5% in the chemotherapy alone group, and 3-year survival was 25.7% in the mistletoe group compared to 14.2% in the chemotherapy alone group.</li> </ul>
Hamrin et al (2018) (106)	Prospective	N: 52 Ca type: Breast Cancer Stage: Not specified	Agent: Not reported Dose: Not reported Route: Not reported Administration: For at least 2 weeks Comparison: Conventional care alone	Conventional care	Immune Response	<ul> <li>Mistletoe group had significantly less CD8 T-cells compared to control (p=0.05), no other immune parameters differed between groups.</li> <li>Anxiety decreased (p=0.04), physical symptoms improved (p=0.05) in the mistletoe group.</li> </ul>
Schad et al (2018) (84)	Retrospective	N: 56 Ca type: Multiple types Stage: I-IV	Agent: Helixor Dose: Not reported Route: Intravenous Administration: Varied Comparison: Monoclonal antibody alone (n = 8), mistletoe alone (n = 12), combined (n = 43)	Most received chemotherapy or supportive therapy	Safety of VAE with monoclonal antibody therapy	<ul> <li>Overall, 34 patients experienced 142 adverse events.</li> <li>Highest incidence of AEs occurred in the monoclonal antibody group (63% of patients) compared to the combination mistletoe group (56% of patients). Five times higher OR of an AE after treatment with mAB compared to mAB plus VAE (95% CI 1.53-16.14).</li> <li>Rates of serious AEs were similar between groups (2% for mistletoe combination group and 3% for monoclonal antibody alone group).</li> </ul>
Thronicke et al (2018) (85)	Retrospective	N: 310 Ca type: Multiple types Stage: 0-IV	Agent: Fraxini, Quercus, Mali Dose: Not reported Route: SC Administration: Median duration was 3.8 months (114 days) Comparison: Targeted therapy alone	Targeted therapy	Safety with targeted therapy	<ul> <li>Mistletoe + targeted therapy, compared to targeted therapy alone, was associated with a significant reduction in overall AE rate (20.1% vs 35%, p=0.04) and a significant reduction in therapy discontinuation rate (30.2% vs 60.5%, p=0.03).</li> <li>Odds ratio of discontinuation of treatment was 0.30 for the mistletoe + conventional care group (p=0.02).</li> </ul>
Oei et al (2019) (91)	Retrospective	N: 106 Ca type: Multiple Cancer Types & Multiple Auto- Immune Diseases Stage:0-IV (most were early stages)	Agent: Abnoba, Iscador and Helixor  Dose: varied, escalating  Route: SC (+/- IV) or IV alone or intratumoral  Administration: SC, 2 or 3 times per week. For IV, the	Most received chemotherapy with IV applications	Safety AEs	<ul> <li>84% of the study population reported 0 adverse events related to mistletoe.</li> <li>15% of patients had 1-3 adverse events related to mistletoe and 1 patient experienced 10.</li> <li>Of the 37 mistletoe related AEs, 20 were expected (local reaction &lt; 5 cm, indurations, local injection site reaction). 17 were considered unexpected.</li> </ul>

			dose and administration were varied  Comparison: None			<ul> <li>No patient had to stop mistletoe therapy.</li> <li>In a subgroup analysis of 30 patients with long-term mistletoe therapy, none experienced a flare up/exacerbation of their auto-immune condition.</li> </ul>
Oei et al (2020) (65)	Retrospective	N: 319 Ca type: Breast cancer Stage:Non- metastatic	Agent: AbnobaViscum, Helixor, Iscador, and Iscucin Dose: Not reported Route: SC and IV Administration: Either alone or with chemotherapy. Duration ≥4 weeks Comparison: Chemotherapy alone, mistletoe alone, combined therapy, or no mistletoe or chemotherapy (control – this group could receive endocrine therapy/immunotherapy)	All patients offered standard oncology therapies	Internal coherence (marker of resilience, optimism, sense of control) (ICS questionnaire) Cancer-related fatigue (EORTC QLQ C30) QOL (EORTC QLQ C30)	<ul> <li>Authors report that patients receiving VAE but no chemotherapy experienced significant beneficial effects on thermo-coherence (p&lt;0.05), affective fatigue (p&lt;0.05), and seven EORTC subscales at 24 months (all p&lt;0.05).</li> <li>Chemo-, immuno- and endocrine therapies had a 17-, 17- and 6-point decline, respectively, for EORTC fatigue (P = 0.0004), whereas the VAE group improved 12 points.</li> <li>VAE group improved in insomnia and physical functioning scores while these scores worsened in conventional care groups (p = 0.009 and p = 0.005, respectively).</li> <li>Caution is advised when reviewing these results given the possibility of selective reporting and questionable statistical analysis. Additionally, note that most positive results were for the VAE-only group not VAE + chemotherapy.</li> </ul>
Thronicke et al (2020) (62)	Retrospective	N: 88 Ca type: pancreatic cancer Stage: IV	Agent: Abnobaviscum, Helixor, and Iscador  Dose: Not reported  Route: Mainly SC. IV and intratumoral was performed in individual cases  Administration: Duration for ≥4 weeks  Comparison: Standard care alone	Standard of care	Cost-effectiveness of VAE Overall survival (OS)	<ul> <li>Median OS was 2.8 months longer in mistletoe group compared to standard care alone (p = 0.008), mean OS was 3.5 months longer in the mistletoe group (no P value provided).</li> <li>The addition of the VAE to standard treatment resulted in 1.16 days and 1.43 days longer for mean hospital stays and mean hospitalization length however the results were not statistically significant (p&gt;0.05).</li> <li>Costs per mean month of OS and per mean hospital stay were lower for VAE + standard care compared to standard treatment, however, there was no statistical analysis for this outcome.</li> </ul>
Thronicke et al (2020) (63)	Retrospective	N: 118 Ca Type: NSCLC Stage: IV	Agent: Abnobaviscum, Helixor, and Iscador  Dose: Not reported  Route: Mainly SC (20 and 2 patients also received IV and intratumoral, respectively)  Administration: Duration for ≥4 weeks  Comparison: Chemotherapy alone	chemotherapy	Cost-effectiveness (CE) of VAE Overall survival (OS)	<ul> <li>VAE + standard care group had longer age-adjusted mean overall survival (OS) than standard care alone group (19.1 months versus 13.4 months, respectively). No statistical analysis was applied to determine significance.</li> <li>Compared to the control group, patients in the VAE group had a lower cost per mean months OS. No statistical analysis was applied to determine significance.</li> </ul>
Thronicke et al (2020) (64)	Retrospective	N: 275 Ca type: NSCLC patients Stage: I -IIIA	Agent: Abnobaviscum, Helixor, and Iscador Dose:	Standard oncological treatment	Overall survival (OS)	- There was no significant difference in OS between the VAE + standard care and standard care alone groups.

			Route: SC route or by off-label IV administration (52.6% of patients) Administration: duration for ≥4 weeks Comparison: Standard oncological treatment alone				
Baek et al (2021) (59)	Retrospective	N: 52 Ca type: rectal adenocarcinoma Stage: II-III	Agent: Abnoba Viscum Q Dose: dose escalation every 3 weeks from 0.02mg to 20mg Route: SC Administration:3X/week for 3 weeks Comparison: neoadjuvant chemoradiotherapy alone	Neoadjuvant chemoradiotherapy	Tumor response	- Tu co sig reg (8) (8) - Ly gr - No	AE group (N=15) compared to a no-VAE group (N=37). Yumor response was significantly better in the VAE group compared to the no-VAE group, meeting statistical significance in pCR rate (53.5% vs 21.6%, p=0.044), tumor egression grade (66.7% vs 32.4%, p=0.024), T downstaging 86.7% vs 43.2%, p=0.004), overall TNM downstaging 86.7% vs 56.8%, p=0.040). Cymphovascular invasion was more common in the no VAE roup (32.4% vs 13.3%, p=0.04). Us significant differences seen in adverse effects, with the nost common toxicity in both groups being stage 1 proctitis.

ADR; adverse drug reaction, AE; adverse event, Ca; cancer, CTCAE; common terminology for adverse events, N; number, QOL; quality of life, SC; subcutaneous, Tx; treatment,

# **Disclaimer**

This monograph provides a summary of available evidence and neither advocates for nor against the use of a particular therapy. Every effort is made to ensure the information included in this monograph is accurate at the time it is published. Prior to using a new therapy or product, always consult a licensed health care provider. The information in this monograph should not be interpreted as medical advice nor should it replace the advice of a qualified health care provider.

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