Nutritional management and prevention of oral mucositis in haematology and oncology cancer patients undergoing antineoplastic treatments

Balma García-Gozalbo and Luis Cabañas-Alite

Abstract: Cancer is a prevalent disease worldwide and treatments such as radiotherapy and chemotherapy sometimes lead to adverse events. Oral mucositis is one of the most disabling and clinical guidelines do not take into account nutritional interventions. The primary endpoint was to gather the evidence about the efficacy of nutritional interventions in the prevention and/or treatment of antineoplastic induced oral mucositis in oncological patients. It was carried out a bibliographic review in PubMed data base by combining MesH terms with boolean operators. Articles were selected based on inclusion and exclusion criteria; 50 final articles were found. Although further evidence is needed, glutamine, honey and vitamins appear to be a good therapeutic option. The rest of the compounds presented controversial or insufficient results to draw conclusions over their utilization as prevention or treatment options. Low evidence is reported about oral mucositis nutritional interventions in spite of being attainable and affordable compounds. Scarce evidence is shown in paediatric patients compared to adults. Developing higher quality studies and combinations with the compounds researched is necessary to create stronger evidence.

Keywords: cancer; oral mucositis; diet therapy

1. Introduction

The World Health Organization (WHO) conceptualizes cancer as a generic term which includes a wide group of diseases that can affect any part of the organism. They are also called “malignant tumours” or “malignant neoplasms”. A defining characteristic of cancer is the rapid division of abnormal cells that extend beyond the common limits and can invade adjacent parts of the body or spread to other organs, a process called metastasis. Metastasis is the main cause of death due to cancer disease [1]. It is indispensable to highlight the word “group” of diseases, considering that they are different entities with common characteristics such as their genetical origin, uncontrolled proliferation, resistance to cell death and capability to invade adjacent tissues or metastasize in distant organs [2]. Based on this definition, we can indicate the global situation on morbidity and mortality on account of cancer. From the data reported by the international agency for cancer research (IARC) in 2018, we can elucidate that the incidence went up to 18.1 million cases and the mortality scaled to 9.6 million. Also, lung, colorectal and mammary cancers are still the most incident of all. Furthermore, women death rates are lower than men rates [3]. When neoplastic processes are diagnosed, they come alongside a treatment. Whether chemotherapy or radiotherapy is used, adverse events are found, being oral mucositis one of the most common among others such as diarrhoea or vomiting [4]. It is estimated that conditioning regimens for stem cell transplantation are the ones that generate the highest rates of oral mucositis incidence, followed by radiotherapy and lastly by chemotherapy [5-9].
All things considered; it is possible now to define oral mucositis more specifically, which is defined as the inflammation of oral mucosa with clinical consequences such as ulcers or erythema secondary to radiotherapeutic or chemotherapeutic treatments [10]. This term differs from the unit "mucositis", which is considered the damage and inflammation of oral, pharyngeal, laryngeal and oesophageal mucosa, together with other areas of the gastrointestinal tract secondary to antineoplastic treatments such as radiotherapy and chemotherapy [8]. Likewise, sometimes the word "stomatitis" is used as a synonym for oral mucositis (OM) even though it is a different entity, because it defines the inflammation in the oral mucosa due to other specific aetiology unrelated to antineoplastic treatments (e.g. infections) [10,11].

Multiple scales are used to assess OM, but none of them is agreed as a standardized scale. The classification divides them into general scales (e.g. WHO scale), scales with multiple variables (e.g. Beck, Eliers and Walsh scales) and the ones specific for the treatment (e.g. Oral Mucositis Rating, National Cancer Institute and Radiation Therapy Oncology Group/European Organization for Research and Treatment of Cancer scales) [12-14].

After OM is evaluated, it is possible to find either mild erythema and burnt mouth sensation or profound mouth ulcers and the inability to eat. Independently of the clinical presentation, its cause relies on a matrix of heterogeneous processes. Not only one physiopathological mechanism is the direct cause for OM but many complex biological routes. OM is typically based on five stages. The first two of them have an immediate appearance after chemotheraphy, radiotherapy or chemoradiotherapy. The first phase is the so-called "initiation" and it is based on the death of basal epithelial cells (as a consequence of the treatment damage to the DNA in cells) along with the secretion of reactive oxygen species (ROS) and endogenous damage-associated pattern molecules (CRAMPs). The last of which, binds to specific receptors and set in motion the second stage. This second step is designated such as "primary damage response” and involves a cascade of biological events which interact with one another and conclude activating various transcription factors (nuclear factor Kappa-B (NF-κB), Wnt, p53, and their associated canonical pathways). For example, NF-κB route activation can be done directly (by the antineoplastic treatments) or indirectly (because of ROS and CRAMPs), evidencing the complexity of OM physiopathology. The activation of transcription factors produces the expression of a large number of genes, being some of them related to the production of molecules (eg, COX-2, inducible NO-synthase, superoxide dismutase...) which take part in the secondary emergence of OM. Other routes directly linked with the onset of this pathology exist, such as the nitrogen metabolism pathway, ceramide and fibrinolysis route, and the stimulation of matrix metalloproteinases (MMPs).

The third stage is called "signal amplification” and occurs when the primary response molecules have positive or negative feedback on the local tissue. Throughout this period there is no visible injury even though the submucous tissues and basal membrane are already damaged. Four to five days after the antineoplastic treatment, the destructive processes of the three first stages triggers before the fourth or "ulceration stage” begins. This phase implies the ulceration of the oral mucosa (transecting the full epithelial thickness), being the patients more prone to infections when this happens. Moreover, ulcers are colonized by oral bacteria worsening and making the initial injury last longer due to infiltrating macrophages generating pro-inflammatory cytokines. The "healing stage" or last phase consists of the remission of the oral cavity injuries spontaneously. This event happens as a result of the activation from signaling molecules (extracellular matrix) which direct the migration, proliferation, and differentiation of the epithelium bordering ulcerative areas [11].

Managing OM is complex. Clinical practice guidelines (CPG) on this topic are scarce and not recent in its vast majority, with exceptions. The Multinational Association of Supportive Care in Cancer and International Society of Oral Oncology (MASCC/ISOO)
has the most recent clinical practice guideline on oral mucositis prevention and treatment topic.

Strategies such as basic oral care are beneficial practices with a low grade of evidence. Therefore, procedures such as a multiagent combination of oral care protocols (a guide about the time, frequency and products that patients with cancer should use every day) are considered beneficial to prevent OM from appearing and others like chlorhexidine rinses are contraindicated to prevent it (grade III evidence), nevertheless saline or bicarbonate sodium rinses, patient education and professional oral care have insufficient evidence to determine whether they positively or negatively impact OM. It is recommended to use benzydamerinses to prevent OM in patients with head and neck cancers undergoing radiotherapy (RT) or chemotherapy (CT) (grade I and II evidence respectively). Other anti-inflammatory drugs were studied and none of them had enough evidence to present a recommendation [8,15,16].

Regarding other drugs, sucralfate is not recommended as prevention or treatment, whereas topical morphine (0,2%) is suggested for OM treatment when it is associated with pain (low evidence grade III) in patients with head and neck cancer undergoing CT and RT. Oral cryotherapy (ice therapy), in contrast, is recommended to prevent OM onset thirty minutes before receiving 5-fluorouracil (5-FU) boluses meanwhile CT, or whether a patient is treated with melphalan in high doses previous to an autologous stem cell transplantation (which means that the donor and receptor are the same human)[8,15,16].

The use of intravenous keratin growth factor (KGF-1) is recommended for the prevention of OM in patients affected by haematologic cancer before an autologous stem cell transplantation with conditioning regimens including high dose CT and total body irradiation (TBI). In this context, neither topical granulocyte-macrophage colony-stimulating factor (GM-CSF) nor parenteral glutamine should be used in the prevention of OM in stem cell transplantation in contrast to what has been said about KFG-1. However, experts suggest using oral glutamine to prevent OM from appearing in patients with head and neck cancer undergoing CT and RT (grade II evidence). Finally, it is considered to use honey to prevent OM in patients with head and neck cancer who are treated with RT or CT and RT [15-16].

The existing clinical practice guidelines about OM don’t or poorly address nutritional approach treatment, leaving unresolved doubts and evidence voids on which interventions should be done. This study pretends to address which nutrients can be administered to prevent and/or treat CT and RT induced OM in cancer patients.

2. Materials and Methods

   Literature Research

   Main aim: To establish the effect of nutritional interventions on the prevention or treatment of oral mucositis in cancer patients undergoing radiotherapy and/or chemotherapy. This paper is based on a bibliographic review of the published information from 2000 to 2021. The literature chosen to be reviewed dates back from October 2020 to April 2021, which includes the latest results of the research strategy used (see supplementary material).

   The bibliographic research was carried out at PubMed and Medline database, finding 252 results.

   Study Eligibility, selection and Data Extraction.

   On one hand, research articles were selected taking into account the following inclusion criteria: 1) original articles, 2) articles published between 2000 and 2021, and 3) those written in English and Spanish. On the other hand, exclusion criteria were: 1) reviews with or without meta-analysis, 2) topics of the article unrelated to the research subject matter, 3) articles whose aim was treating animals as a veterinary care strategy, and 4) qualitative research articles.

   The screening process can be observed in Figure 1.
The review process was completed by two reviewers independently and blindfolded, in other words, without knowing the other reviewer decisions, the scientific journal, the reference or the author.

As shown in Figure 1, 46 articles were excluded after the first screening process made by reading the title and abstract of each article. Out of these 46 articles, 9 of them were excluded due to their language, 1 because of duplication, and 36 for being reviews. In this article, 208 references were selected beforehand to be submitted to a second screening process which consisted of a full reading and an analysis of its relevance to be included in the review. Those references that were not considered useful for this study were excluded (158).

3. Results

Following the literature research, 254 articles were identified. After excluding 204 of them, 50 articles that met the inclusion criteria remained and were included in the final review. Figure 1 further outlines the selection process used. A narrative analysis of the studies revealed two types of population (paediatric and adults), and three key themes emerged: Glutamine, Honey and Other Dietary Components or Prevention Methods. The results are classified according to these themes. Articles included in the final review can also be seen in Table 1.

3.1. Summary of the Studies Included

**Paedriatic population**

Of the final articles included, six of them approached the prevention and/or treatment of OM in paediatric patients with cancer. Ages evaluated in those articles went from one to nineteen years old, varying in the different scientific articles. Also, all the articles majorly approached haematological neoplasms, even though some included solid tumours in their sample (in lessening proportion). Four of the studies were based on
clinical trials, whereas the other two were based on a cohort study and a case series study. The four trials studied different compounds.

**Glutamine**

From the reviewed articles, 18 articles retrieved were related to different preparations of glutamine for the prevention and treatment of OM in adults with cancer. All the studies were performed in population that ranged from 19 to 70 years old. The most frequent ages groups were those ranging 30 and 50 years old. The overall articles consisted of 15 clinical trials, 1 cohort study and 2 laboratory studies (one in vivo and the other in vitro and in vivo).

Regarding clinical trials, it can be stated that their nature is heterogeneous. More specifically due to the fact that different levels of methodological blindness exist (double and triple blindness, open trials, preliminary, etc.), and there are differences in the treated disease, the used therapy (CT, RT or CRT), even OM grading scale.

**Honey**

Another compound studied as a possible prevention and/or treatment for OM is honey. Having retrieved after a preliminary research, 8 articles about it. Out of these, 7 articles were designed as a research with different types of blinded clinical trials and 1 of them was a laboratory study (in vitro and in vivo).

**Other Dietary Components or Prevention Methods**

Vitamins were another group that was analysed in different studies, (7 articles found).

Liquorice was studied in 3 clinical trials, establishing different results.

Cryotherapy was examined in 2 articles in which was determined a reduction of OM incidence and severity.

It must be highlighted that there are also two compounds analysed (date palm pollen and polideoxyribonucleotides) in the present study which weren't possible to compare with the existent scientific literature since they have only been researched in 1 article. However, their results were positive, being in need of further evidence to provide new approaches to the prevention and treatment of OM induced by antineoplastic therapies.
<table>
<thead>
<tr>
<th>Author (year)</th>
<th>N</th>
<th>Study design</th>
<th>Objective</th>
<th>Intervention</th>
<th>Time (months)</th>
<th>Conclusion</th>
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</thead>
</table>
| Huang et al. (2019) | 59 | RCT Phase III, double-blind | To evaluate if oral glutamine prevents acute toxicities (OM and dermatitis) secondary to the treatment with RT in patients with head and neck cancers. | - Glutamine (TG).  
- Control: maltodextrin (CG)  
NCT CTCAE 4.03 version | 19 | CG developed more OM (grades 2-4) than TG; nevertheless, the efficacy of the treatment with oral glutamine is not meaningful after RT in head and neck cancers. |
| Tsujimoto et al. (2015) | 40 | RCT, double-blind | To evaluate if oral glutamine reduces mucositis severity induced by CRT in patients with head and neck cancer. | - Glutamine (TG).  
- Placebo (PG).  
NCT CTCAE 3.0 version | 36 | Oral glutamine reduced OM severity produced by CT in head and neck cancer patients with a maximum mean grade of OM lower for TG than to PG, a duration without meaningful difference between both groups and a less duration of artificial nutrition required in TG. |
| Tanaka et al. (2016) | 30 | RCT Phase II | To assess if glutamine and the combination of glutamine and elemental diet reduce the incidence of CT induced OM in patients with oesophageal cancers. | - Placebo (PG)  
- Glutamine (TG)  
- Glutamine + elemental diet (CTG).  
NCT CTCAE 3.0 version | 36 | Without differences between PG and TG |
| Pachón et al. (2018) | 262 | Prospective cohort study | To evaluate if oral glutamine prevents mucositis induced by oncological therapies (CRT or RT) in patients with head and neck cancer. | - Oral glutamine (TG).  
- Placebo (PG).  
RTOG/EORTC (MO andoesophagitis)  
| Chang et al. (2019) | 60 | RCT Triple-blind | To measure the impact of oral glutamine as a supplement on the prevention of oesophagitis induced by CRT in patients with advanced non-small cell lung cancer (stages III-IV). | - Placebo (PG)  
- Oral glutamine (TG).  
ARIE: acute radiation induced oesophagitis | 12 | TG had less severe ARIE than PG. Also a reduction of the incidence in the weight loss in TG. |
<p>| Anturlikaret al. (2019) | 20 | In vitro and in vivo study | To measure the security and efficacy of “HTOR-091516” (Tumeric, Triphala and honey) as a treatment of OM induced by 5-FU | IN VITRO: gingival human fibroblast, mouse connective tissue and human oral reconstructed | 0.5 (14 days) | The average weight loss in TG was lower, there also was less mortality and a reduction in OM grade (WHO scale). The product is proposed to prevent OM. |
| Cho et al. (2019) | 91 | CT | To assess the effect of glutamine enriched parenteral nutrition (PN) on weight, infections, complications (mucositis, neutropenia and graft versus host disease) and mortality in patients who undergone hematopoietic stem cell transplantation. | epidermis culture. Later treatment with “HTOR-091516” and MTT test (cytotoxicity) + TNF-α inhibition test (inflammation)+ test INVITTOX SKINETHIC™ (irritation). IN VIVO: two groups of rats (control [PG] and treatment with “HTOR-091516” [TG]). In TG: drops on the induced ulcer in every animal during their treatment with CT. | 48 | Without significative association in the case of OM duration. It was noted a descent in 100-days mortality for TG. |
| Shumsky et al. (2019) | 15 | Pilot RCT | To evaluate the efficacy of Oncoxin (ONCX) in oncologic patients with OM who undergo CT, RT or both. |  | 0.6 (20 days) | The was found a lower OM grade in OG (after 7 days of treatment and towards the end of their treatment). |</p>
<table>
<thead>
<tr>
<th>Study</th>
<th>Population</th>
<th>Design</th>
<th>Objective</th>
<th>Intervention</th>
<th>Duration</th>
<th>Results</th>
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</thead>
<tbody>
<tr>
<td>Widjaja et al. (2020)</td>
<td>48 Double-blind RCT</td>
<td></td>
<td>To measure if oral glutamine prevents OM during CT (metotrexate [MTX] in paediatric patients with acute lymphoblastic leukaemia (ALL))</td>
<td>- Placebo group (PG). - Oral glutamine group (TG) WHO Scale</td>
<td>0.5 (14 days)</td>
<td>There was a descent on the incidence and severity of OM in TG after CT.</td>
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<tr>
<td>Harada et al. (2018)</td>
<td>50 In vitro and in vivo study</td>
<td></td>
<td>To appraise the efficacy of Elental© (dietetic liquid formulation enriched with amino acids, which is a source of L-glutamine) as 5-FU (CT) induced OM and dermatitis treatment.</td>
<td>IN VIVO: - Saline solution (PG). - Dextrin group (DG). - Elental group (EG).</td>
<td>0.25 (8 days)</td>
<td>In vivo, OM healing faster than in EG</td>
</tr>
<tr>
<td>Oosterom et al. (2019)</td>
<td>99 (A) y 81 (B) Cohort study</td>
<td></td>
<td>A. To study the prevalence of vitamin D deficiency in paediatric patients</td>
<td>- Vitamin D levels before and after MTX therapy. - After MTX, classification in two groups depending on OM grade: ≥3 or ≤3 NCI CTCAE 3.0 version</td>
<td>2004 – 2012</td>
<td>There was no association between basal vitamin D levels and MTX induced OM, but low levels of vitamin D during MTX therapy were found to be related to severe OM.</td>
</tr>
<tr>
<td>Sun et al (2019)</td>
<td>100 Double-blind RCT</td>
<td></td>
<td>To examine the effects of a group B multivitamin complex combined with GeneTime© (human recombinant growth factor) on the treatment of OM in patients with head and neck cancer undergoing RT.</td>
<td>- Control group with vitamin B complex. - Observational group with (OG) vitamin B complex + GeneTime© RTOG scale.</td>
<td>12</td>
<td>Less severity, affected area and healing time of the OM ulcers in OG.</td>
</tr>
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</table>
Nejatinamini et al. (2018) 28 Cohort study  To evaluate the changes in vitamin status during the treatment of head and neck cancers related to body composition, inflammation and mucositis.

- Dietetic intake measurement (3 days).
- Vitamin levels (vitamins D, E, B9 and B12)
- Basal (before treatment) and 6 to 8 weeks after RT treatment (with or without CT) C-reactive protein (CRP) measurement.
- C-reactive protein (CRP) baseline before and after 6 to 8 weeks of RT treatment, either with or without QT after treatment.

1 - 1.5 Higher rates of OM were observed related to less vitamin D, B12, E and B9 intake and lower blood levels of vitamin A and D.

Tanaka et al. (2018) 19 RCT To measure the intake of Elental® during two cycles of CT and to determine the incidence of OM in patients with oesophageal cancers treated with CT who completed their intake and those who did not completed it.

- Elental® group (CG).
- Elental® group with uncompleted treatment (UG).

CTCAE 3.0 version

2 (56 days) Less severity of OM in CG during CT with the use of Elental®.

Pathak et al. 56 RCT To assess the efficacy and role of oral - Control group (CG).

1.75 (49 days) TG had less hospitalizations due to OM and disphagia.
<table>
<thead>
<tr>
<th>Year</th>
<th>Study Design</th>
<th>Participants</th>
<th>Interventions</th>
<th>Comparison</th>
<th>Findings</th>
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<tbody>
<tr>
<td>2019</td>
<td>glutamine in the treatment of OM and dysphagia induced by chemoradiotherapy (CRT) in patients with oropharynx and larynx carcinoma.</td>
<td>- Glutamine group (TG). NCI CTCAE 4.03 version</td>
<td>More incidence and severity of OM in CG.</td>
<td></td>
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<tr>
<td>Mamgain et al. (2020)</td>
<td>RCT</td>
<td>To evaluate the efficacy of an ayurvedic preparation (based on <em>Glycyrrhiza glabra</em>) in order to decrease the severity of mucositis in patients with head and neck cancers who receive chemoradiotherapy (CRT).</td>
<td>Comparison between basal and post-RT characteristics: &lt;1. Conventional OM treatment (antiacids and anaesthetics) (CTG) &lt;2. Conventional treatment and ayurvedic preparation (ATG). &lt;3. Honey and conventional treatment (HTG).</td>
<td>24 Less severity, pain and onset time of mucositis, both in ATG and HTG, but especially in ATG.</td>
<td></td>
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<tr>
<td>Harada et al. (2019)</td>
<td>Open RCT</td>
<td>To evaluate changes in OM (injuries’ size, pain and redness + CRP in plasma) in patients with oral squamous cell carcinoma undergoing CRT or RT with Elental® administration.</td>
<td>- Elental Group (EG). - Non-Elental Group as the control (CG). CTCAE 4.0 version</td>
<td>24 In EG, milder OM development in CRT; no difference in RT.</td>
<td></td>
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<td>Rao et al. (2017)</td>
<td>Blinded</td>
<td>To evaluate whether honey causes</td>
<td>- Povidone-iodine</td>
<td>6 Lesser OM incidence and severity in HG. The implication on</td>
<td></td>
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<tr>
<td>Study</td>
<td>Design</td>
<td>Population</td>
<td>Intervention</td>
<td>Outcomes</td>
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</table>
| Branda et al. (2004) | 68 Pilot cohort study | To study the influence of **B12 vitamin, folate and dietetic supplements** on CT-induced toxicity in breast cancer patients. | - Blood samples (B12, B9 and neutrophils) before/after the first CT cycle.  
- Questionnaires about supplement usage.  
- OM grading (author-modified CTCAE).  
- 68 subjects + historical controls. | No evidence on the influence was found. |
| Okada et al. (2017) | 20 Pilot single-centre RCT | To evaluate the influence of **Elental®** on CT-induced OM and diarrhoeas in patients with oesophageal cancer. | - IG: use of Elental®.  
- CG: no Elental used.  
- Questionnaires and clinical examination  
- CTCAE 4.0 version | Less severe OM incidence in IG. |
| De Sousa et al. (2018) | 40 In vivo study | To evaluate the effects of **glycine** in the expression of collagen and platelet and epidermal growth factors (PDGE, EGF) in a OM murine model. | - Control group (CG).  
- Intervention group (IG: glycine supplementation). | Positive effects in IG, with a better recovering rate (collagen increase and growth factors reduction). |
<table>
<thead>
<tr>
<th>Reference</th>
<th>Group Size</th>
<th>Study Design</th>
<th>Objective</th>
<th>Description</th>
<th>Duration</th>
<th>Results</th>
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<tr>
<td>Nihei et al. (2018)</td>
<td>67</td>
<td>Single-centre RCT</td>
<td>To evaluate the efficiency and safety of L-Glutamine sodium azulene sulphonate in the treatment of CT-induced OM in patients with colorectal and breast cancer.</td>
<td>- Intervention group (IG). - Control group (CG: standard oral hygiene). CTCAE 4.0 version NRS pain scale</td>
<td>24</td>
<td>Lesser OM severity in IG. No significant differences were found as for the incidence.</td>
</tr>
<tr>
<td>Chattopadhyay et al. (2014)</td>
<td>70</td>
<td>Single-centre RCT</td>
<td>To evaluate the influence of oral glutamine on RT-induced OM in patients with head and neck cancer.</td>
<td>- Intervention group with oral glutamine (IG). - No placebo control group (CG). WHO scale.</td>
<td>8</td>
<td>Lower incidence, severity and duration of RT-induced OM in IG. Menor incidencia, severidad y duración de MO inducida por RT en GI.</td>
</tr>
<tr>
<td>Üçüncü et al. (2006)</td>
<td>35</td>
<td>Laboratory CT (rats)</td>
<td>To determine the preventive effect of Vit E (VE) and L-Carnitine (LC), alone or in combination, on OM and myelosuppression by RT.</td>
<td>- 5 groups: 1) No RT (control: saline + simulated radiation). 2) RT. 3) RT+VE. 4) RT+LC. 5) RT+VE+LC. - OM measurement scale: Parkins et al. - Clinical/histopathological follow-up: 4 days pre-RT – 10 days post-RT.</td>
<td></td>
<td>VE and LC proved to be radioprotective agents on their own and not combined together, with lower severity and longer time to histological appearance of OM. Good tolerance and no adverse effects.</td>
</tr>
</tbody>
</table>
Amanat et al. (2017) 82 Single-centre RCT To assess the effect of honey on clinical grades of OM.  
- Honey group (HG).  
- Control saline group (CG).  
RTOG scale.  
12 Lower incidence and severity of OM in HG, during the RT.

Podlesko et al. (2018) 3 Case series To evaluate the effects of topical application of deoxyribonucleic acid on 3 OM (moderate-severe) cases in patients with head and neck cancer.  
Oral spray of polydeoxyribonucleotide (PDRN) as treatment.  
1 Increased relief and remission of OM as a matter of time, without interruption of the treatment or opioids intake.

Perrone et al. (2017) 73 CT To analyse the influence of dietary supplementation with whey protein concentrate (WPC) on the incidence of OM in patients undergoing HSCT.  
- WPC Group (WG).  
- Historical controls (CG).  
- WG was sub-stratified into: consumed <80% PWC (WG1) or≥80% (WG2) of the offered dose.  
WHO and CTCAE 4.0 version  
Not specified No significant differences between WG and CG in what incidence, duration and severity of MO concerns. However, in WG2, shorter duration and lower incidence of severe OM was found.

Ogata et al. (2017) 22 Pilot prospective To evaluate the preventive effects of Elental® on CT-induced OM in  
- 22 patients with Elental (1 group).  
36 Significantly reduced CT (5-FU)-induced OM grade.
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>Description</th>
<th>Outcome</th>
<th>Control Group</th>
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</thead>
<tbody>
<tr>
<td>Al Jouni et al. (2017)</td>
<td>Open RCT</td>
<td>To evaluate the effects of honey on grade 3-4 OM, reduction of bacterial/fungal infections, duration of OM episodes and body weight in paediatric leukaemia patients undergoing CT or RT.</td>
<td>12 Significant reduction in severity and pain in HG. Significant improvement in weight and time to OM onset in HG.</td>
<td>Control group (CG) with Lidocaine, Mycostatin, Daktarin and oral cleaning. Experimental group (HG) with same routine as CG + honey (4-6 times/day). WHO scale</td>
</tr>
<tr>
<td>Lopez-Vaquero et al. (2017)</td>
<td>Phase II double-blind RCT</td>
<td>Evaluates whether glutamine is effective in reducing the incidence and severity of mucositis and dermatitis induced by RT or CRT in patients with head and neck cancer.</td>
<td>6 Incidence and severity with no significant differences between groups.</td>
<td>L-Glutamine group (TG). Placebo group with malto-dextrine (PG). WHO scale</td>
</tr>
<tr>
<td>Howlader et al. (2019)</td>
<td>RCT (single-blinded)</td>
<td>To assess whether honey improves mucositis injuries and the life quality of patients with RT/CT-induced OM (for head and neck cancer).</td>
<td>From CT start – 4 weeks after RT. Less OM and associated symptoms induced by RT in HG. Shorter time towards the recovery of a regular life quality.</td>
<td>Treatment group (HG) with honey (both, rinsed and ingested honey). Control group (CG) with saline solution. WHO scale</td>
</tr>
<tr>
<td>Elsass, F.T.</td>
<td>Case series</td>
<td>The aim was to improve OM (oral application of honey).</td>
<td>- Application of honey - Shorter healing time with lower pain rate in all cases.</td>
<td>-</td>
</tr>
<tr>
<td>Year</td>
<td>Study Design</td>
<td>Study Description</td>
<td>Interventions</td>
<td>Duration</td>
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<td>------------------------------------------------------------------------------------</td>
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<tr>
<td>(2017)</td>
<td>(3 analyzed)</td>
<td>comfort and feeding) with standard oral care and the use of Leptospermum honey in paediatric oncology patients after proven CT.</td>
<td>on the buccal surface with a cotton swab, 3 times/day. Then spat or sucked out.</td>
<td></td>
</tr>
</tbody>
</table>
| Elkerm y Tawashi (2014) | 20 Pilot study | To evaluate whether date palm pollen (DPP) can be effective in the prevention and treatment of RT and CT-induced OM in patients with head and neck cancer. | - DPP Group (one daily suspension).  
- Control group (CG) (antifungal, rebamipide and oral analgesia).  
OMAS score and visual analogue scale for mouth pain and dysphagia. | 1.5 (6 weeks) | Significant reduction in incidence, severity and pain in OM and dysphagia in DPP. |
| Raeessi et al. (2014) | 61 Single-centre double-blind RCT | To evaluate the effects of coffee + honey in the treatment of OM by CT and to compare them with the effects of steroids. | 3 grupos:  
- Betamethasone group (EG).  
- Honey group (HG).  
- Honey + coffee group (HCG).  
WHO scale | 36 2011-2013 | Significant reduction in the severity of MO in all three groups. |
<p>| Baydar et al. (2005) | 99 CT | To research the effects of local cryotherapy on the prevention of CT (5-FU)-induced OM. | - Intervention Group (IG): CT courses with local cryotherapy (ice in the mouth during the CT course up to 10 days) | Not specified | 5-FU-induced OM incidence lower in GI (OR=11.5). |</p>
<table>
<thead>
<tr>
<th>Study</th>
<th>Participants</th>
<th>Design</th>
<th>Objective</th>
<th>Intervention Details</th>
<th>Results/Findings</th>
</tr>
</thead>
</table>
| Peterson et al. (2007) | 305          | Phase III, double-blind RCT | To observe the efficacy of Saforis® in the prevention and treatment of OM caused by CT treatment in breast cancer. | - Control Group (CG): no-cryotherapy courses.  
- Saforis group (SG).  
- Placebo group (PG) (with subsequent cross-linking).  
WHO scale (ulcer grading scale); OMAS scale (ulceration measurement) | Not specified. Lower severity and incidence rate in SG.                                                                 |
| Fogh et al. (2016)    | 119          | Multicentric phase II RCT | To evaluate the effect of Manuka honey (liquid and tablets) in the prevention of RT-induced oesophagitis in lung cancer patients. | - G1: Manuka honey (liquid).  
- G2: Manuka honey (tablets).  
- G3: control, standard care.  
Measurements: odynophagia (NRPS scale), pain, opioid use, dysphagia, weight loss, quality of life and nutritional status. | 12 There were no significant differences for groups G1, G2 or G3, so the use of honey did not prove to be superior to standard health care. |
| Samdariya et al. (2015)| 69           | Open RCT               | To study the intake of honey in pain relief caused by RT-induced OM in patients with head and neck cancer. | A. Gargle with soda and benzidamine (PG).  
B. Gargle with soda + benzidamine + honey. | Nov. 2011 – Jan. 2013. Slightly greater relief in HG during the entire follow-up (3 months), with a significant reduction in the severity of OM-associated pain and fewer treatment interruptions. |
<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>Phase</th>
<th>Purpose</th>
<th>Intervention</th>
<th>Duration (days)</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Matsuda et al. (2015)</td>
<td>Double-blind phase III RCT.</td>
<td>90</td>
<td>To research whether TJ-14 (Hangeshashinto) prevents and/or controls CT-induced OM in patients with colorectal cancer.</td>
<td>TJ-14 treatment group (IG). - Placebo group (PG). WHO scale.</td>
<td>0.5 (14 days).</td>
<td>Significant reduction in the duration of severe OM, with no effect on the severity or incidence of OM itself.</td>
</tr>
<tr>
<td>Jayachandran y Balaji (2012)</td>
<td>RCT</td>
<td>60</td>
<td>To evaluate the effect of natural honey and benzidamine hydrochloride on the development and severity of RT-associated OM in patients with oral cancers.</td>
<td>3 grupos (oral rinses): A. Honey (HG). B. Benzidamine hydrochloride (BG). C. Saline 0.9% (control group, CG).</td>
<td>6</td>
<td>Lower severity and earlier healing of OM not significant in HG.</td>
</tr>
<tr>
<td>Study</td>
<td>Total N</td>
<td>Study Design</td>
<td>Objective</td>
<td>Findings</td>
<td>Duration</td>
<td>Notes</td>
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<tr>
<td>Sugita et al. (2012)</td>
<td>118</td>
<td>Retrospective CT</td>
<td>To ascertain the effects of folinic acid administration (systemic and rinsed) on the incidence of OM and acute graft-versus-host disease (GVHD) after GVHD prophylaxis with MTX in patients undergoing HSCT.</td>
<td>Systemic folic acid was administered to patients at increased risk of developing OM (n=29).</td>
<td>48</td>
<td>It could be useful in reducing the incidence of severe OM, both in systemic use and rinsed.</td>
</tr>
<tr>
<td>Sorensen et al. (2008)</td>
<td>206</td>
<td>Double-blind RCT</td>
<td>To evaluate prevention of OM using chlorhexidine compared to cryotherapy during 5-FU CT in gastrointestinal cancer.</td>
<td>3 groups:</td>
<td>3</td>
<td>Higher severity rate of OM in PG and lower in CG, with a lower incidence and duration in IG and CG than in PG, therefore prophylaxis seems effective in both IG and CG.</td>
</tr>
<tr>
<td>Das et al. (2011)</td>
<td>52</td>
<td>RCT</td>
<td>Observe protective/healing effect against RT and CT effects (OM, skin reaction, xerostomia or voice changes) when using Glycyrrhiza glabra.</td>
<td>4 groups:</td>
<td>1.75 (7 weeks)</td>
<td>Lower incidence and severity in GLHG and LHG compared to CG, but similar to HG.</td>
</tr>
<tr>
<td>Thornley et al. (2004)</td>
<td>37</td>
<td>CT</td>
<td>To determine the feasibility and potential efficacy of a fixed combination of agents in reducing RRT (regimen-related toxicity) in children undergoing HSCT.</td>
<td>Combination group of ursodeoxycholic acid (UDCA), vitamin E, folinic acid and titrated parenteral nutrition (IG).</td>
<td>36</td>
<td>Significant decrease in the incidence and severity of OM in IG.</td>
</tr>
<tr>
<td>Study</td>
<td>Participants</td>
<td>Design</td>
<td>Objective</td>
<td>Intervention</td>
<td>Controls</td>
<td>Study Duration</td>
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<tr>
<td>Iyama et al. (2014)</td>
<td>44 RCT</td>
<td>To research whether supplementation with GFO (Glutamine, Fibre, Oligosaccharides) decreases the severity of mucosal injury post-HSCT.</td>
<td>- Historical controls group (1995-2000) (PG).</td>
<td>- GFO group (IG). - Control (CG).</td>
<td>CTCAE 4.0 version</td>
<td>36</td>
</tr>
<tr>
<td>Takano et al. (2015)</td>
<td>96 In vitro study</td>
<td>To investigate whether γ-tocotrienol (vitamin E) can enhance survival of oral human keratinocytes (RT7) against 5-FU-induced cell toxicity.</td>
<td>- RT7 cells were treated with 5-FU and γ-tocotrienol. - 4 groups: A: γ-tocotrienol. B: 5-FU. C: γ-tocotrienol + 5-FU. D: Control de 5-FU + N-acetylcysteine.</td>
<td>-</td>
<td>-</td>
<td>In C there was a significant inhibition of ROS production induced by 5-FU.</td>
</tr>
<tr>
<td>Agha-Hosseini et al. (2021)</td>
<td>59 Triple-blinded RCT</td>
<td>To evaluate whether a vitamin E, hyaluronic acid and triamcinolone mouthwash is effective in the treatment of radiotherapy-induced OM grades 3-4.</td>
<td>- Group with vitamin E+hyaluronic acid+triamcinolone rinses (IG). - Group with triamcinolone rinses (CG).</td>
<td>WHO scale</td>
<td>4 weeks</td>
<td>Significant reduction in the severity of RT-induced OM in IG overtime.</td>
</tr>
<tr>
<td>Study</td>
<td>Sample Size</td>
<td>Design</td>
<td>Objective</td>
<td>TG Intervention</td>
<td>PG Intervention</td>
<td>WHO Scale</td>
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<tr>
<td>Yeshurun et al. (2020)</td>
<td>52</td>
<td>Multicentric double blind RCT</td>
<td>To determine whether folinic acid (FA) reduces methotrexate (MTX)-induced toxicity in patients undergoing myeloablative conditioning (CM) for allogeneic haematopoietic cell transplantation, who have as well received MTX prophylaxis for graft-versus-host disease.</td>
<td>TG with FA</td>
<td>PG with placebo.</td>
<td>17 (4.5-50)</td>
</tr>
<tr>
<td>Pattanakitsakul et al. (2020)</td>
<td>30</td>
<td>Preliminary and unicentric quasi-randomized trial</td>
<td>To examine the protective effect of vitamin A supplementation against mucosal damage of the gastrointestinal tract after CT in paediatric patients undergoing HSCT. As a secondary objective, to assess the occurrence of OM.</td>
<td>TG with single dose (200000UI) of vitamin A.</td>
<td>PG without vitamin A.</td>
<td>12</td>
</tr>
</tbody>
</table>
4. Discussion

**PEDIATRIC EVIDENCE.**

Al Jaouni et al. (2017), throughout an open clinical trial, determined that honey diminished the incidence of the worst oral mucositis grades and delayed its onset. It was also encountered that honey helped to decrease pain, infections and hospital stay. Furthermore, increase patients weight compared with the control group. In contrast, Widjaja et al. (2020), with their double-blind randomized controlled trial showed less incidence of OM in their treatment group with glutamine, proving other results such as a decrease in the severity of OM due to the glutamine and less time of treatments and sanitary costs.

Alternatively, Thornley et al. (2004) demonstrated an OM incidence and severity reduction along with less regimen-related toxicity (especially in high-risk patients) thanks to a preparation made of ursodeoxycholic acid, vitamin E, folic acid and parenteral nutrition as prevention. In contrast, Pattanakitsakul et al. (2020) proved in their preliminary quasi-randomized trial that vitamin A did not prevent OM.

Oosterom et al. (2019) in their cohort study evidenced that basal vitamin D levels were not related with MTX induced OM, but it was found an association (OR=1.26) between the reduction in vitamin D levels during MTX treatment with severe OM (grade 3 or more). Finally, a case series (3) reported the benefits of honey as OM treatment in patients undergoing chemotherapy.

Sung et al. (2015) carried out a clinical practice guideline where they collected all the available evidence about glutamine (and other treatments) to prevent OM in paediatric patients undergoing a conditioning regimen previous to a hematopoietic stem cell transplantation, and they determined that there was no consistency on the reduction of OM in more than one study. The only retrieved study about glutamine in paediatric patients on this paper reveals a change, given that it could be determined that this clinical trial is consistent evidence of the effectiveness of glutamine to decrease the incidence of OM, highlighting that more studies are needed to consider it as a recommendation (Widjaja et al., 2020).

In addition, the same Sung et al. (2015) established in their review that topical vitamin E did not demonstrate to reduce OM effectively. In the present study, it was not possible to compare the retrieved article with Sung et al. (2015) review since it was applied a combination of agents. Nevertheless, it must be emphasised the huge lack of studies with vitamins, it is necessary to identify their effect and implement recommendations based on sufficient data (Thornley et al., 2004).

The cohort study with vitamin D was not possible to compare with others because of the inexistence of more papers, corroborating the previous paragraph about the lack of evidence in this subject (Oosterom et al., 2019).

Friend et al. (2018) clarified in their review that honey could be effective in the treatment and prevention of OM in paediatric patients in limited resources areas, but determines that there is no available evidence that compares if honey is even or more effective than the established treatments. Evidence found in this study supports this affirmation since the open clinical trial shows the efficacy of honey in OM, but there are no comparative studies between honey and other treatments. Also, case series don't have that strong evidence to be compared with clinical trials or to state conclusions if more evidence doesn't exist (Al Jaouni et al., 2017).

**GLUTAMINE**

Regarding adults and the use of glutamine, Nihei et al. (2018), Chattopadhyay et al. (2014) and López-Vaquero et al. (2017) in their scientific studies didn't show a reduction in the total incidence of OM. In contrast, Huang et al. (2019) and Tanaka et al. (2016) evidenced a reduction in the total incidence of OM; this matches the contribution of a cohort study made by Pachón et al. (2018), except for the fact that they didn't report less incidence in the severity of OM (grade ≥2). The trials of Huang et al. (2019) and López-Vaquero et al. (2017) got the same result on this matter, contrasting with the re-
sults of Tsujimoto et al. (2015), Tanaka et al. (2016), Nihei et al. (2018) and Chattopadhyay et al. (2014) who observed less severity of OM in patients treated with glutamine.

Concerning the time of onset of OM, only three papers addressed this topic, and two of them determined an absence of delay in the onset of OM, contrarily to what Chattopadhyay et al. (2014) study addressed (Tsujimoto et al., 2015; López-Vaquero et al., 2017). As for the duration of OM, it was studied in four trials, proven in two of them a reduction in the duration of the most severe OM (Chattopadhyay et al. 2014; Iyama et al. 2014).

Other nutritional factors, such as weight, were studied in five scientific articles, showing a tendency in weight loss slightly minor in those groups with glutamine. Also, Tanaka et al. (2016) proved the maintenance of weight in cancer patients when it was mixed glutamine and elemental diet (Iyama et al., 2014; Chang et al., 2019; Cho et al., 2019; Pathak et al., 2019).

Quality of life (QoL) was evaluated by López-Vaquero et al. (2017) through an adapted questionnaire for this topic without obtaining a link between the use of glutamine and OM patients’ quality of life. Other factors such as pain and the use of analgesia (3 articles), dysphagia (1 article) and odynophagia (1 article) were studied. The results provided determined that the treatment with glutamine improved dysphagia, odynophagia and pain, showing two of them a decrease in analgesia use too (Tsujimoto et al., 2015; Nihei et al. (2018); Pachón et al., 2018; Pathak et al., 2019).

Four of the trials delved into a product called Elental®, a liquid dietetic formula enriched with amino acids and a source of L-glutamine. The results showed at large a reduction in the severity of OM in groups treated with Elental®, highlighting the investigation of Harada et al. (2019), who proved a decrease in the administered analgesia and CRP levels (4-6 weeks of CRT). These results concur with the ones obtained by Harada et al. (2018) in their in vivo study in which rats treated with Elental® healed more rapidly of OM ulcers (Ogata et al., 2016; Okada et al. 2017; Tanaka et al., 2018).

About its combination with other products, Peterson et al. (2007) analyzed a compound called Saforis® (an oral formulation that increases the availability of glutamine in the oral cavity), which proved to decrease the severity of antineoplastic induced OM, with no other conclusive results to report. Finally, Bateman et al. (2013) used a combination of WHEY protein, fatty acid and glutamine, showing that this doesn't protect against OM in rats. Nevertheless, Anderson and Lalla (2020) performed a review where they suggested glutamine rinses (liquid formula) in patients with head and neck cancers undergoing CRT since they found that it decreases the severity and duration of OM and oesophagitis. This is comparable to five of the found studies in the bibliographic search, even though all of them study oral glutamine to rinse and afterwards swallowing, revealing the same benefits reported in the commented review. Oral glutamine might be beneficial for both topical and by ingestion, but it should be further studied since the body of evidence must be larger to recommend rinses or the intake of glutamine (Chattopadhyay et al., 2014;Tsujimoto et al., 2015; López-Vaquero et al., 2017; Pachón et al., 2018; Huang et al., 2019).

In addition, recently Shuai et al. (2020) published a meta-analysis where affirmed that oral glutamine could have no clinical benefits in the prevention and/or treatment in CRT or RT induced OM in patients with head and neck cancers. Therefore, glutamine should be studied further in order to obtain more consistent results.

**HONEY**

Severity is the most studied aspect of honey use. It has been proven that honey is effective in the eventual reduction of the severity of OM, as Howlader et al. (2019), Rao et al. (2017) and Amanat et al. (2017) point out in their scientific publications.

Along the same lines, Raeessi et al. (2014) indicated that the efficacy of the combination between honey and coffee to decrease the severity of OM exceed the same capacity of honey by itself and steroids. On the other hand, Fogh et al. (2016) delved into the severity of oesophagitis and they did not find a link between honey and the decrease of OM...
stage. This same study does not highlight the association between honey and a reduction of late forms of odynophagia (it doesn't occur in the same way in precocious forms).

Rao et al. (2017) approaches the incidence of OM and proves the benefits of honey in the prevention and delay of the onset of OM. Also, they showed a reduction in weight loss in their treatment group with this compound. Jayachandran y Balaji (2012) also observed a delay in the onset of OM in patients treated with honey in comparison to those who were treated with benzydamine; in fact, no effects were found on the duration.

Regarding other parameters analysed, only Howlader’s et al. (2019) publication assesses the influence of honey on the quality of life (QoL) during OM, being observed a late improvement of OM because of this. The trial performed by Samdariya et al. (2015) showed a consistent decrease in the severity of the pain, resulting in fewer discontinuances in RT.

As a final remark, Anturlilkar et al. (2019) in their in vitro and in vivo study with a combination of turmeric, Triphala and honey (HTOR-091516) showed positive results such as low product toxicity, inflammation inhibition (TNF-α) and a protector effect in CT induced OM.

Münstedt y Männle (2019) in their review evaluating the use of honey as OM treatment showed a decrease in the severity of the ulcers which harmonize with the scientific articles found in the present paper and in the systematic review and meta-analysis made by Tian et al. (2020).

Other aspects mentioned by Tian et al. (2020) include the indication of a potential decrease in the incidence of OM during the treatment with honey, however, from the present study, only one study proves significant prevention.

There exist other interesting aspects such as OM pain reduction and the inhibition of the inflammatory pathways (in vitro study with “HTOR-091516”) which are not comparable with the mentioned reviews, since there is no evidence to suggest or recommend the use of honey in this aspects. Even though honey could be useful to reduce mucositis severity, the rest of the aspects should be studied more in-depth to provide reliable data and determine suggestions and recommendations on this matter.

VITAMINS AND AMINO ACIDS.

The most important results of the three retrieved clinical trials showed a reduction in the severity of OM in patients treated with a combination of GeneTime®, group B vitamin complex, Oncoxin® (combination of vitamin C, B6 and amino acids) and vitamin B9, proving a reduction in mucositis incidence too. Nevertheless, Branda et al. (2004) in their cohort study didn’t find a link between the quantity of vitamins B12, B9 and multivitamins supplements with this severity (Sugita et al., 2012; Shumsky et al., 2019; Sun et al., 2019).

Specifically, Oncoxin® was related to a weight increase and a normal intake too. On the one hand, the combination of GeneTime® and vitamin B proved less pain and quicker healing of OM ulcers. Finally, the treatment with vitamin B9 coincide with the effect of Oncoxin® in the improvement of the intake capacity, but it didn’t show a reduction in parenteral nutrition use of time; it did show a reduction of opioid use (Sugita et al., 2012; Shumsky et al., 2019; Sun et al., 2019).

On the other hand, Nejatinamini et al. (2018) showed that the decrease in the blood levels of vitamins A and D during RT are related to OM. In the same way, an in vitro study determined relevant information since it showed that vitamin E (γ-tocotrienol) in CT fosters the survival of oral human keratinocytes through the inhibition of reactive oxygen species (ROS), probably by the suppression of Nrf2 route (Takano et al., 2015).

Recently, a clinical trial published by Agha-Hosseini et al. (2021) with a rinse made of vitamin E, hyaluronic acid and triamcinolone proved to be useful in the treatment of OM. Similarly, another scientific group did a clinical trial with folic acid for the prevention of OM. However, their results did not prove to be very successful (Yeshurun et al. 2020).
Related to the use of complete protein, the study of Perrone et al. (2017) proved that there was not a link between the intake of concentrated WHEY protein and the incidence, severity and duration of OM. However, the individuals who took more quantity of WHEY protein presented less severity and duration of OM compared to those who took less quantity of protein. Finally, De Sousa’s et al. (2018) in vivo study indicated that glycine promoted major and better tissue restructuring.

Regarding non-combined amino acids, a clinical trial and an in vivo laboratory study are highlighted. The rest of the studies related to amino acids (3 in total) appear in combination with other nutritional compounds and have already been mentioned.

Yarom et al. (2019) considered that vitamin B9 could not be effective in the prevention of OM in patients who receive therapy for a posterior hematopoietic stem cell transplantation, highlighting a cohort study (which is also analyzed in this study) that proves the efficacy in the reduction of both total incidence and OM severity in patients undergoing CT for a posterior stem cell transplantation. There exist few studies related to vitamin B9 to determine some evidence about it.

Evidence void also occurs when it is studied the treatment of OM with vitamins E and D since both have some studies (5 about vitamin E and 1 about vitamin D) which determine different effects about OM and cannot be compared between them due to the characteristics used to analyze each study (they use different laboratories, cohorts, clinical trials...). Therefore, supporting what Yarom et al. (2019) mention in their review, it is not possible to determine consistent evidence for the use of vitamins.

In this section, when it comes to analyze the use of amino acids, it is highlighted that there are not reviews about the use of Glicine and concentrated WHEY protein. As a result, there is a lack of evidence in this study (2 articles, one of each of the topics) to determine whether they could be useful in treating OM.

**GLYCYRRHIZA GLABRA.**

On the one hand, the study of Mamgain et al. (2020) and Das et al. (2011) found both a reduction in incidence and severity in those groups treated with liquorice (Glycyrrhiza glabra), with more effect than those treated with honey in the same trial. Contrastingly, Matsuda et al. (2015) proved that the use of Hangeshashinto (mixed with 7 medicinal plants, Glycyrrhiza glabra among them) does not affect the incidence and stage of OM, but it does affect in the duration of severe OM.

Finally, highlight that Das et al. (2011) also found fewer interruptions in the treatment and persistence of xerostomia despite liquorice administration.

OM therapy with liquorice is novel and not so much researched. In the present study, there were only the articles analyzed about this compound, making it insufficient to determine clear evidence or preventive and/or healing effect in OM. Richard (2021) proves in their review that glycyrrhizin and glycyrrhetinic acids have anti-inflammatory properties, inhibiting specifically interleukin secretion (IL-6, IL-2, IL-12...), the expression of TNF-α and cytokine cascade, among others.

Even though this literature review does not focus on OM specifically, attention could be drawn towards this matter, since physiopathologically a lot of inflammatory routes are involved in the production of oral ulcers, and compounds such as liquorice (with antiinflammatory properties) could give results in the inhibition of inflammatory routes if further research is carried out in the future.

**OTHERS.**

Elkerm y Tawashi (2014) delved into the part of date pal pollen in OM treatment through a preliminary study. Such research showed a reduction in the incidence and severity of RT induced OM. It was also proven a reduction in the pain and dysphagia during OM.

Secondly, deoxyribonucleic acid was studied by Podlesko et al. (2018), using it as a topical spray in a case series. The results showed pain relief and quicker healing on OM ulcers in two cases and a deterioration in the third one.
López-González et al. (2021) proves the efficacy to prevent and treat OM through oral cryotherapy. This evidence was also found in several clinical practice guidelines, highlighting its best utility in the prevention of OM in patients treated with CT. The present study cannot compare the results of the given evidence due to the scarceness of the analysis (2 articles), but it does agree with the results found in the evidence retrieved from other reviews and clinical practice guidelines (Peterson et al., 2011; Lalla et al., 2014; Elad et al., 2020).

**STRENGTHS**

The performance of this paper allows the gathering of information related to a not so studied topic, opening at the same time future doors to investigations about this subject and evidencing the need for more scientific literature about this pathological entity, prevalent between oncological patients.

In the same way, carrying out this review helps to prove which is the best available evidence on OM nutritional treatment (either prophylactic or for a treatment itself). Likewise, it facilitates a guide for health professionals involved in the patient care process such as dietitian-nutritionists, nurses and doctors.

**LIMITATIONS** (Weaknesses)

Some methodologic limitations must be highlighted in the present study. On the one hand, the available time to carry out this review was highly limited (from October to February), affecting the extension of the bibliographic search. On the other hand, a context limitation was present since it was noted the lack of consensus in the use of OM grading scales in the different scientific articles, making it impossible to compare the articles between them even though they assessed the same nutritional compounds.

Lastly, nutritional interventions in this entity aren’t studied as much as they must be studied, therefore, there is not enough scientific literature to affirm solid conclusions, requiring more support with further research articles.

5. Conclusions

In paediatric ages, positive results are found for the treatment and/or prevention of OM, but it is needed more investigation since it does exist an evidence void for the use of all the studied nutritional compounds.

Regarding to the adults population, glutamine and honey could be the most useful treatment for OM, but more evidence is needed to confirm a reduction in the severity. In vitamins and amino acids (different from glutamine), little evidence exists. It seems like vitamins could serve as a treatment, but there is scarce evidence and generally, the investigated formulations don’t only include vitamins, making it difficult to prove their effects on their own.

It is not possible to present the usefulness of Glycyrrhiza glabra, date palm pollen and polydeoxyribonucleotides in the prevention and/or treatment of OM in oncological patients due to a lack of evidence. Moreover, cryotherapy was also analyzed insufficiently in this study, contrasting with what was found in the scientific literature.

In general, very few scientific productions exist about the nutritional approach of CT and/or RT induced OM in cancer patients even though the high interest of the study for being an alternative treatment is highly attainable and easily available.

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**References**


