Regulatory aspects and evidences of melatonin use for sleep disorders and insomnia: an integrative review

Aspectos regulatórios e evidências do uso de melatonina em distúrbio do sono e insônia: uma revisão integrativa

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ABSTRACT

Background: Insomnia is a sleep disorder characterized by difficulty of falling asleep or maintaining sleep, which affects different age groups. Currently, melatonin is used as a therapeutic treatment in cases of insomnia in children, adults, and elderly people. Objective: To evaluate the effectiveness of melatonin in sleep disorders, its dosage, potential adverse effects, as well as labeling laws and regulations in Brazil. Methods: This integrative review was carried out using the Cochrane Library, Medline (Pubmed), and Science Direct databases. Twenty-five articles and three documents available on the Brazilian Society of Endocrinology and Metabolism (SBEM) and National Health Surveillance Agency (ANVISA) websites published between 2015 and 2020 were selected to be evaluated in full. Results: It was found that in most of the selected articles the use of melatonin reduces sleep latency. The effective melatonin doses varied according to each age group, from 0.5 to 3 mg in children, 3 to 5 mg in adolescents, 1 to 5 mg in adults, and 1 to 6 mg in elderly people. Side effects are mild when taking usual doses. In Brazil, no registered drug and current regulation on the use and marketing of melatonin has been identified. Conclusion: The use of melatonin is an alternative therapy that can be used for sleeping disorders. According to the evidences found, it did not demonstrate toxicity or severe side effects, nor dependence even when administered at high doses, suggesting that it is a safe medication to treat patients of different ages suffering from sleeping disorders.

Keywords: Melatonin; Sleep Wake Disorders; Pineal Gland.

RESUMO

Antecedentes: Insônia é um distúrbio do sono caracterizado por dificuldade de iniciar e manter o sono, afetando diferentes faixas etárias. Atualmente, a melatonina é utilizada no tratamento de insônia em crianças, adultos e idosos. Objetivo: Avaliar a eficácia da melatonina nos distúrbios do sono, posologia e potenciais efeitos adversos, bem como a regulamentação vigente no Brasil. Métodos: Trata-se de uma revisão integrativa, os artigos foram identificados nas bases de dados Cochrane Library, Medline (Pubmed) e Science Direct, totalizando 25 artigos, e foram selecionados três materiais disponíveis no site da Sociedade Brasileira de Endocrinologia e Metabologia e Agência Nacional de Vigilância Sanitária, publicados entre 2015 e 2020. Resultados: Verificou-se na maioria dos artigos selecionados que a melatonina reduz a latência do sono. Quanto as dosagens de melatonina identificou-se variação em cada faixa etária, para crianças de 0,5 a 3mg; adolescentes de 3 a 5mg; adultos de 1 a 5mg e idosos 1 mg a 6 mg demonstraram serem eficazes. Em doses habituais os efeitos colaterais são leves. No Brasil, não foi identificado medicamento registrado e regulamentação vigente sobre o uso e comercialização de melatonina. Conclusão: A utilização da melatonina é uma alternativa que pode ser utilizada em distúrbios do sono. De acordo com as evidências encontradas, não demonstrou toxicidade ou efeitos colaterais severos, nem dependência mesmo em doses elevadas, sendo, portanto, segura para tratamento de pacientes desde crianças a idosos que sofrem de distúrbios do sono.

Palavras-chave: Melatonina; Transtornos do Sono-Vigília; Glândula pineal.
INTRODUCTION

Insomnia is a sleep disorder characterized by difficulty in falling asleep or maintaining sleep. It is defined as the persistent difficulty in falling and staying asleep, problems with sleep duration and maintenance and sleep quality. Insomnia occurs in different age groups and can last for weeks, months, or longer periods. In these cases, it is considered as a chronic disease when it remains for three months or more and affects the individual’s occupational performance and daily routine. The prevalence of insomnia varies between 10 to 20% in the general population, and approximately 50% of these people live with this condition in a chronic way. In China, the prevalence of insomnia is 15%, followed by Spain 21.1% and Brazil, reaching above 30% of the population. The gold standard for sleep assessment, this study aimed to describe the objective prevalence of insomnia in the São Paulo, Brazil, Epidemiologic Sleep Study cohort of 1,101 adults (20-80 years old).

Non-pharmacological treatments can help the patient in improving symptoms, including sleep hygiene, stimulus control, relaxation techniques, among other methods. Pharmacological treatments used to treat insomnia include selective agonists of the γ-aminobutyric acid type A (GABA-A) receptor, sedative antidepressants, melatonin and melatoninergic agonist, sedative antipsychotics, benzodiazepines, anticonvulsants, antihistamines, and herbal medicines such as Valeriana officinalis. Widely used, the minor tranquilizers benzodiazepines can cause tolerance, dependence, and withdrawal syndrome, as well as being considered inappropriate for elderly people due to its high risk of accidents, including falls and concomitant fractures.

Commonly known as melatonin, n-acetyl-5-methoxytryptamine is a neurohormone, a small lipophilic molecule produced by the pineal gland. Among its functions, the chronobiotic effect is associated with the regulation of the endogenous clock in relation to the photoperiod. A physiological function of endogenous melatonin is to reinforce the behavior related to darkness. Its production increases about two hours before bed. In addition, during the night, melatonin is responsible for transmitting information to the brain and other organs of the central nervous system about the duration of sleep, reducing the watch signal, as well as promoting fatigue and inducing sleep. Li et al. Embase, Cochrane Library, ClinicalTrials.gov, and Web of Science reviewing exogenous melatonin as a treatment for secondary sleep disorders suggested that exogenous melatonin improves sleep quality, reduces onset latency, and increases total sleep time.

Classified as a non-prescription supplement in the United States of America, melatonin is widely used as a natural product, among all age groups, including children. Research conducted with 31 supplements in Guelph, Ontario, Canada has shown low quality of melatonin formulations, with high concentration variability between samples and batches and the presence of serotonin in 8 of the evaluated supplements. In Brazil, the number of medical prescriptions for the use of melatonin to treat insomnia has increased, however their commercialization is available only in compounding pharmacies. Currently, only one supplier of pharmaceutical products is authorized to distribute melatonin in Brazil, and solely to compounding pharmacies. Although the use of melatonin for the treatment of insomnia has expanded in recent years, information about its potential adverse effects and drug interactions are still limited. The National Health Surveillance Agency (ANVISA) newsletter (2019) reports that marketed medications must have proof of safety, efficacy, and quality in Brazil, but due to an injunction that allows the sale of raw materials, melatonin is available over-the-counter. Moreover, the Brazilian Sleep Society considers that the use of melatonin to treat circadian rhythm disorders is already established, but the results for insomnia are not consistent, despite some positive results in specific populations and a good tolerability and safety profile, with few side effects.

These results demonstrate the need for greater control over the production and marketing of melatonin supplements. It is necessary to build and develop a more effective health care and identify the best practices for health professional bodies, especially prescribers and pharmacists who work directly in the prescription and provision of this medication to patients. In this perspective, this study aimed to evaluate the effectiveness of melatonin for sleep disorders and melatonin associated symptoms, dosage, potential adverse effects, as well as the law regulations of the drug in Brazil.

METHODS

This article is an integrative review in which the six methodological steps described by Mendes, Silveira and Galvão were followed: (i) identification of the theme and selection of the research question to carry out the integrative review; (ii) establishment of criteria for inclusion and exclusion of studies; (iii) definition of the information to be extracted from the selected studies; (iv) evaluation of studies included in the integrative review; (v) interpretation of results; and (vi) presentation of the review.

Data collection was performed using Cochrane Library, Medline (Pubmed), and Science Direct databases. To perform the searches, Health Sciences Descriptors (Decs) were used in Portuguese and English, and the articles were selected according to the objective of the project and including the following key words: melatonin, pineal gland, sleep disorders, and insomnia. In addition, for the regulatory aspects, the Brazilian Society of Endocrinology and Metabolology (SBEM) and the ANVISA websites were accessed using the descriptor “melatonin”. Fifty-six documents were found using the
SBEM and two documents were selected to be evaluated in full. Thirty-four documents were identified using the ANVISA website and one document was selected to be assessed in full.

The article search using the databases was based on the following criteria: articles published between 2015 and 2020, in Portuguese or English. Articles that were not in the defined languages, did not have any descriptors of interest, or that were outside the defined publication period were excluded. Reviews, meta-analysis, cohort studies, and randomized clinical trials were selected.

For the selection of articles, an initial search based on title and abstract was carried out in the database. Then, a new selection was made with the reading of the articles in full, according to the Figure 1.

| 1st Phase: Guiding Question |
|----------------------------|
| "What are the scientific evidences regarding the use of Melatonin for sleeping disorders?" |

| 2nd Phase: Data collection |
|----------------------------|
| Definition of databases and search for articles |
| PUBMED | COCHRANE LIBRARY | SCIENCE DIRECT |
| 179 | 16 | 178 |

| 3rd Phase: Articles that contained descriptors or search expressions in the title or abstract |
|------------------------------------------------------------------------------------------|
| 28 | 4 | 09 |

| 4th Phase: Articles that suggested addressing the objective |
|----------------------------------------------------------|
| 18 | 2 | 5 |

| 5th Phase: Articles that answered the guiding question |
|-------------------------------------------------------|
| Total of selected articles = 25 |

Figure 1. Summary of the article selection process.

RESULTS

The final selection consisted of 25 articles and three documents identified in the Brazilian websites. Table 1 summarizes the characteristics of the articles according to authors, year of publication, and database of publication.

No articles were identified reporting the current regulations of melatonin in Brazil. The only information found was that the commercialization of melatonin is authorized only for compounding pharmacies, and must be purchased from the Active Pharmaceutica supplier. Similarly, no drug is registered with melatonin as the active ingredient. However, in Europe, a medicine registered under the trade name of Circadin is available in the market.

Table 2 shows the characteristics of the selected articles according to the study design, population, dosage, intervention evaluation, therapeutic and adverse effects of melatonin. Of the 25 studies found, 22 showed positive results regarding the use of melatonin in insomnia disorders and three studies showed ineffective or inconclusive results.

DISCUSSION

The term circadian is derived from the Latin term *circa diem*, which means around one day. Circadian rhythms are endogenous oscillations that occur over a 24-hour period. In humans, this cycle lasts an average of 24.2 hours with individual variation of 23.7 to 25.3 hours. The waking and sleeping process is strongly influenced by the circadian system.

Currently approved by the Food and Drug Administration (FDA) for the treatment of sleep disorders in elderly people, melatonin has been used for delayed insomnia, sleep-wake cycle with periods shorter than 24 hours, sleep correction in the elderly, as an adjuvant in the treatment of autism spectrum disorder, attention deficit hyperactivity syndrome, migraine, anesthesia, metabolic diseases, and polycystic ovary syndrome. The synthetically produced melatonin can be administered exogenously and there are immediate and sustained release formulations available on the market.

Malow and collaborators report that melatonin is effective for sleep disorders in children and adolescents.
Table 1. Distribution of references included in the study, according to the authors, year of publication, database and journal.

| Authors | Year | Data base | Periodic |
|---------|------|-----------|----------|
| Hajak & Zisapel | 2015 | PUBMED | International clinical psychopharmacology |
| Culpepper & Wingertzahn | 2015 | PUBMED | Prim Care Companion CNS Disord |
| Wright et al. | 2015 | SCIENCE DIRECT | Drugs & Aging |
| Foley & Steel | 2015 | SCIENCE DIRECT | Complementary Therapies in Medicine |
| Williams et al. | 2016 | PUBMED | Pharmacotherapy |
| Mcleery & Sharpley | 2016 | COCHRANE LIBRARY | Cochrane Database of Systematic Reviews |
| Cardinali et al. | 2016 | SCIENCE DIRECT | Pharmacological Research |
| Chang et al. | 2016 | PUBMED | JAMA pediatrics. |
| Hohl et al. | 2016 | SITE SBEM | SITE SBEM |
| Sociedade Brasileira de Endocrinologia e Metabologia | 2016 | SITE SBEM | SITE SBEM |
| Auld et al. | 2017 | PUBMED | Sleep Medicine Reviews |
| Madsen et al. | 2017 | PUBMED | Trials journal |
| Riemann et al. | 2017 | PUBMED | European Sleep Research Society |
| Abdelgadir et al. | 2018 | PUBMED | Archives of Disease Childhood |
| Maras et al. | 2018 | PUBMED | Journal Of Child And Adolescent Psychopharmacology |
| Quera-Salva & Clausrat | 2018 | PUBMED | Encephale |
| Zwart et al. | 2018 | PUBMED | Healthcare |
| Myers et al. | 2018 | PUBMED | Journal of clinical sleep medicine |
| Sletten et al. | 2018 | PUBMED | Public Library of Science |
| Lewis et al. | 2018 | COCHRANE LIBRARY | Cochrane Database of Systematic Reviews |
| Anvisa | 2019 | SITE ANVISA | SITE ANVISA |
| Besag et al. | 2019 | PUBMED | CNS Drugs |
| Schroder et al. | 2019 | PUBMED | Journal of autism and developmental disorders |
| Lemoine; Bablon & Silva | 2019 | PUBMED | Complementary therapies in medicine |
| Seiden & Shah | 2019 | PUBMED | Prim Care Companion CNS Disord |
| Li et al. | 2019 | PUBMED | Frontiers in neuroendocrinology |
| Malow et al. | 2020 | SCIENCE DIRECT | Journal of the American Academy of Child & Adolescent Psychiatry |
| Low, Choo & Tan | 2020 | SCIENCE DIRECT | Journal of Psychiatric Research |

(2-17.5 years) with autism spectrum disorder and insomnia. According to the authors, there was an improvement in sleep quality with no changes in weight, height, body mass index, and pubertal status, and no evidence of developmental delays. Nunes et al. recommend doses of 0.5 to 3 mg for children and 3 to 5 mg for adolescents. Chang et al. suggest that melatonin supplementation is a relatively safe and effective way to improve sleep-onset in addition to decreasing the severity of symptoms of atopic dermatitis in children, due to melatonin immunomodulatory, anti-inflammatory, and antioxidant effects, thus improving the skin and helping to maintain the epidermal barrier.

The review demonstrated that the use of melatonin is effective to treat primary and secondary insomnia at different stages of life, from children and teenagers to adults and the elderly. There was also a great diversity in the variables investigated, especially regarding the dose, time of use, and sleep outcomes. Despite of some biases, most studies demonstrated that the use of melatonin for sleep disorders is efficient and safe.

Table 3 shows the dosages of melatonin indicated for each age group according to the publications found.

As for the use in elderly people, Culpepper et al. report that prolonged-release melatonin formulations are effective in symptoms associated with sleep-onset, however the effects may be limited to individuals over 55 years old who suffer from insomnia. Quera-Salva et al. evaluated a dose of 2 mg of melatonin administered once a day for three months. The authors found that Circadin® is well tolerated, has no rebound, withdrawal or hangover effects, in addition to not causing drug interactions with antihypertensive, antidiabetic, hypolipidemic or anti-inflammatory drugs, which are the drugs most used by elderly people.

No evidence was found that the use of melatonin (up to 10 mg) assists in sleep disorders in patients with moderate to severe dementia due to Alzheimer’s disease. The doses...
| Authors                  | Study type                   | Population | Aims                                                                 | Posology                   | Evaluation of the intervention                                                                 | Effects of melatonin in insomnia |
|-------------------------|------------------------------|------------|----------------------------------------------------------------------|----------------------------|-------------------------------------------------------------------------------------------------|----------------------------------|
| Hajak & Zisapel          | Prospective cohort study     | 597 patients (210 men and 387 women) | To investigate the effects of interruption, withdrawal and rebound from Circadin®. | 2 mg, 2 hours before sleep for 3 weeks | Sleep quality was classified as 1 = very good, 2 = good, 3 = regular, 4 = poor and 5 = very poor. Morning alertness was classified as 1 = fully alert, 2 = alert, 3 = regular, 4 = tired and 5 = very tired. Improvement or worsening of sleep quality or morning alertness was defined as a change (decrease) of at least one point from the baseline. | 75% of patients reported that sleep quality and morning alertness had improved to at least regular to good or very good. No serious adverse effects have been reported. |
| Culpepper & Wingertzahn  | Systematic review            | 1344 patients aged 55 years old     | To investigate the level of evidence that supports the use of over-the-counter agents: diphenhydramine, doxylamine, melatonin and valerian for occasional sleep disorders or insomnia. | 5 mg daily for 8 weeks, 2 mg daily for 3 weeks | Primary: sleep latency actigraphy, sleep time, awakenings, sleep efficiency. Secondary: sleep diary, awakenings, alertness. | It appears to be effective for symptoms associated with onset of sleep and shows a favorable tolerability profile, but the effects may be limited to individuals over 55 years old with insomnia. |
| Wright et al.            | Systematic review and meta-analysis | 322 patients with a mean age of 64 years old | To determine the effect of melatonin compared to placebo discontinuation of benzodiazepines, in addition to determining the effect of melatonin on sleep quality in this population. | 2 to 5 mg once a day before bed 4 -18 weeks | Pittsburgh sleep quality score. | The effect of melatonin on sleep quality was inconsistent. |
| Foley & Steel            | Systematic review            | Patients from 1 to 8 years old; 50 to 85 years old with Alzheimer’s, 16-25 years old with chronic sleep problems, 3 to 16 years old with autism and sleep disorders | To explore the available clinical evidence on the safety of oral use of melatonin. | 2 to 10 mg, Period: 3 to 5 years old | Questionnaire applied weekly. Hematology, electrolytes, urinalysis, physical examination. Holter monitoring 24 hours by electrocardiogram. Selection of cognitive, psychomotor, dexterity and memory recall tasks. Simulated driving task, sedation scale and radioimmunoassay. | Supplementation appears to be relatively safe. Adverse events are generally minor. |
| Williams et al.          | Review                       | Patients older than 55 years old    | To compare the pharmacokinetic and pharmacodynamic properties of agomelatine, prolonged release melatonin and ramelteon, to examine the impact of pharmacological properties on clinical efficacy. | 2 mg day, 1 - 2 hours before bed for up to 13 weeks | Sleep latency, sleep onset time, number and duration of naps. | Adverse reactions: headache, nasopharyngitis, back pain, arthralgia, and potential for hepatic metabolism interaction with other drugs. Pharmacokinetic parameters: Absolute bioavailability 15%, T 1/2, average 3.5-4.0 hours, T max, interval, hours 0.75–3.0. In vitro protein binding: ~ 60%, mainly for albumin, α1-acid glycoprotein and high-density lipoprotein. |
| Mccleery & Sharpley       | Systematic review            | 222 patients                           | To evaluate common adverse effects, compared to placebo for sleep disorders in people with dementia. | Up to 10 mg 8 to 10 weeks | Total night sleep time, proportion of day sleep to night sleep. | There is no evidence that it helped with sleep disorders in patients with moderate to severe dementia due to Alzheimer’s disease. |
| Authors          | Study type                              | Population                          | Aims                                                                 | Posology                  | Evaluation of the intervention                                                                 | Effects of melatonin in insomnia                                                                 |
|------------------|-----------------------------------------|--------------------------------------|----------------------------------------------------------------------|---------------------------|-----------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------|
| Cardinali et al. | Review                                   | Patients aged 20 to 90 years old     | To discuss the available data on the effectiveness of melatonin to reduce chronic use of benzodiazepine drugs in patients with insomnia. | 1 mg, 2 mg, 3 mg, 5 mg, 10 mg 1 day - 18 months | Daily records of sleep and quality of wakefulness filled in by patients, and polysomnography. | It has a very safe profile, is well tolerated and, in some studies, has been administered to patients in very large doses and for long periods, without any potential for drug abuse. Several studies have found that more than 50% of patients treated with benzodiazepines discontinue use after treatment with melatonin. Melatonin may become the therapy of choice to reduce dependence on benzodiazepine medications. |
| Chang et al.     | Randomized, double-blind, placebo-controlled clinical trial | 73 children and adolescents of 1 to 18 years old with attention deficit. | To evaluate the effectiveness of melatonin supplementation to improve sleep disorders and disease severity in children with Atopic Dermatitis. | 3 mg or placebo daily for 4 weeks | Actigraphy, subjective change in sleep and dermatitis, sleep variables measured by polysomnography, nocturnal urinary levels of 6-sulatoxymelatonin and serum immunoglobulin levels. | Melatonin supplementation is a safe and effective way to improve sleep onset latency and disease severity in children with Atopic Dermatitis. |
| Auld et al.      | Systematic review and meta-analysis      | 1500 patients of 18 and 80 years old | To assess the evidence base for the therapeutic effects of exogenous melatonin in treating primary sleep disorders | 0.1 a 10 mg for five weeks | Sleep parameters: primary insomnia, delayed sleep phase syndrome, non 24 hours sleep wake syndrome in blind patients and REM-behaviour disorder | Reduction in the time to fall a sleep between the effect on sleep onset latency for melatonin in patients with primary insomnia. Treating delayed sleep phase syndrome demonstrated an overall significant improvement in sleep onset latency compared with placebo. |
| Madsen et al.   | Randomized, double-blind, placebo-controlled clinical trial | 240 patients | To investigate whether prophylactic treatment with melatonin has a preventive effect on depression, depressive and anxiety symptoms, sleep and circadian disorders after Acute Coronary Syndrome. | 25 mg for 12 weeks | Actigraphy, Pittsburgh sleep diary, pain, anxiety, fatigue and general well-being measured by visual scales. | It may have advantages due to its low toxicity, as well as its proven anxiolytic and hypnotic effects. |
| Riemann et al.   | Clinical Guideline                      | Different populations with insomnia, 4099 patients | To provide clinical recommendations for the management of adult patients with insomnia. | 0.3 - 40 mg 12 weeks | Sleep parameters: objective and subjective, number of awakenings, sleep efficiency; sleep onset latency; total sleep time, awakenings after sleep onset. | Reduces sleep onset latency, improves sleep quality. However, the effects were small from a clinical point of view. It has been regarded as a safe medicine. |
| Abdelgadir et al. | Systematic review and meta-analysis      | 682 children aged <18 years with neurodevelopmental disorders | To determine the efficacy and safety of melatonin as therapy for sleep problems in children with neurodevelopmental disorders | 0.1 - 12 mg 1 a 13 weeks | Sleep parameters: total sleep time, sleep onset latency, frequency of nocturnal awakening, adverse events and child’s behaviour | Melatonin improves total sleep time and sleep onset latency in children with neurodevelopmental disorders, with few adverse events related. |
| Authors          | Study type                      | Population                                      | Aims                                                                 | Posology          | Evaluation of the intervention                                                                 | Effects of melatonin in insomnia |
|------------------|---------------------------------|-------------------------------------------------|----------------------------------------------------------------------|-------------------|-------------------------------------------------------------------------------------------------|----------------------------------|
| Maras et al.32    | A prospective double-blind randomized placebo-controlled | 95 patients aged between 2 to 17.5 years old with Autism Spectrum Disorder. | To describe long-term efficacy and safety of pediatric-appropriate prolonged-release melatonin at the optimal daily dose and impact of the treatment on caregivers’ sleep, daytime sleepiness, and quality of life. | 2.5 a 10mg for 13, 39 and 52 weeks | Sleep parameters: Sleep and Nap Diary, Composite Sleep Disturbance Index, caregiver’s Pittsburgh Sleep Quality Index, Epworth Sleepiness Scale, and quality of life. | After 52 weeks of continuous treatment subjects slept 62.06 minutes longer, fell asleep 48.6 minutes faster (p <0.001), had 89.1 minutes longer uninterrupted sleep episodes, less nightly awakenings and better sleep quality compared with baseline and placebo-randomized group. The drug was generally safe; most frequent treatment-related adverse events were fatigue and mood swings. |
| Quera-Salva & Claustra33 | Review                         | Insomnia patients aged 55 years old or older. | To provide data on the physiological and pharmacological effects of melatonin related to sleep. | 2 mg for 3 months | Sleep quality, sleep latency assessed by the falling asleep score, performance the next morning assessed by the behavior score upon awakening and improvement in quality of life. | Benefits: improvement in sleep quality and latency, alertness the next day, and quality of life. Absence of rebound, abstinence or hangover effect, and without safety concerns in concomitant therapy with antihypertensive, antidiabetic, lipid-lowering or anti-inflammatory drugs. |
| Zwat et al.34     | Cohort study                    | 69 children                                      | To assess the discontinuity of melatonin in therapy, the time and the real quality of sleep. To investigate the occurrence of adverse events and the reasons for discontinuing the use of melatonin. | 0.5 - 5 mg        | Pittsburgh Sleep Quality Index, subjective sleep quality, sleep duration, usual sleep efficiency, sleep disorders, use of sleeping medications and daytime dysfunction. Each item is weighted on a scale of 0 to 3. | Long-term therapy appeared to be safe after an average of 71 years of treatment. The results of this study indicate that approximately 75% of children with chronic insomnia in early sleep treated with melatonin will have normal sleep quality without medication ten years later. |
| Myers et al.35    | Randomized, double-blind, placebo-controlled clinical trial. | 13 patients aged between 2 to 50 years old.     | To investigate the efficacy and safety of melatonin in the treatment of sleep disorders in patients with Dravet’s Syndrome. | 6 mg 30 minutes before bed for 2 weeks | Actigraphy, average sleep latency, average sleep efficiency, frequency of seizures, caregiver’s impression of clinical change and adverse events. | No significant adverse events were reported by caregivers. Although the double-blind study found no significant difference in total sleep time, the overall impression from the parent’s report was that melatonin is very beneficial in some patients. |
| Sletten et al.36  | Randomized, double-blind, placebo-controlled clinical trial. | 116 patients with an average age of 29 years old of both genders. | To test the effectiveness of melatonin in patients with Delayed Sleep-Wake Phase Disorder (DSWPD) | 0.5 mg for 4 weeks | Sleep onset time, sleep efficiency. | Short-term and casual administration is an effective and safe treatment for patients with DSWPD with confirmed circadian misalignment, resulting in improvements in objective and subjective quality of sleep, daytime function, and severity of clinical symptoms. |
| Lewis et al.37     | Systematic review               | 151 patients, with 16 admitted to the intensive care unit (ICU). | To assess whether the quantity and quality of sleep can be improved by administering melatonin to adults in the ICU. To assess whether melatonin improves physical and psychological results. | 3 and 10 mg, Orally or enterally for a minimum of two days or until discharge from the ICU. | Sleep quantity and quality, measured by polysomnography, actigraphy, bispectral index or electroencephalogram. | Insufficient evidence was found to determine whether administration would improve the quality and quantity of sleep in ICU patients. Sparse data and differences were found in the study methodology, in the ICU sedation protocols, and in the methods used to measure and report sleep. |
| Authors                        | Study type                  | Population                      | Aims                                                                 | Posology                                      | Evaluation of the intervention                                                                                           | Effects of melatonin in insomnia                                                                 |
|-------------------------------|-----------------------------|---------------------------------|----------------------------------------------------------------------|-----------------------------------------------|--------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|
| Besag et al.39                | Systematic review           | 1625 participants between 1 and 93 years old | To assess the evidence for adverse events associated with short-term and longer-term treatment for sleep disorders | 0.15 - 12 mg/day for 1 to 29 weeks           | To evaluate the occurrence of adverse effects with the use of melatonin in the short and long term.                      | There were no serious adverse effects associated with the use of melatonin in the short term, the most frequent adverse effects were daytime sleepiness and headache. There is scarcity of data with long-term use. |
| Schroder et al.39             | Double-blind placebo-controlled clinical trial | 125 individuals aged 2 to 17.5 years old with Autism Spectrum Disorder or Smith-Magenis syndrome | To evaluate the efficacy and safety of prolonged release melatonin mini-pills in improving the duration and onset of sleep in patients with Autism Spectrum Disorder or Smith-Magenis syndrome. | 3-5 mg for 13 weeks. | World Health Organization Welfare Index, Pittsburgh Sleep Quality Index and the Epworth Sleepiness Scale. | The treatment improved externalizing behaviors (hyperactivity-inattention and conduct) in children and adolescents. The prolonged-release formulation was effective in improving sleep initiation and maintenance, as well as being safe. |
| Lemoine; Bablon & Silva40     | Prospective Study           | 40 participants between 20 and 75 years old | To investigate the effect of a combination of melatonin, vitamin B6 and medicinal plants in patients with mild to moderate sleep disorders. | 1 mg of melatonin; 0.42mg of vitamin B6; 8.4 mg of California Poppy; 150 mg of passion fruit extract; 240 mg of lemon balm for 14 days. | Total sleep duration, sleep onset latency, number of nightmares per night and number of daytime naps and their duration, daytime fatigue was also subjectively assessed by participants in their sleep diary. | There was an improvement in sleep quality, latency of sleep onset, total sleep duration and daytime parameters related to sleep. No serious adverse events have been reported. |
| Seiden & Shah41               | Randomized clinical trial   | 10 patients, 4 men, 6 women aged between 18 and 40 years old | To evaluate the pharmacokinetic and safety profile of a new melatonin formulation in comparison with the melatonin of immediate release (IR-melatonin). | 5 mg. The subjects were confined to the clinic, included daytime dosing, blood collection and exposure to standard ambient light, to prevent significant endogenous production of melatonin. | For each individual, the melatonin plasma concentration time data was used to calculate the following pharmacokinetic parameters: maximum concentration, maximum time and threshold (time above the target threshold concentration). | The formulation demonstrated rapid release and then continuous release and absorption of melatonin for up to 7 hours, making it a significant advance in the pharmacokinetic release profile of exogenous melatonin delivery and therefore an important potential consideration as a therapy for sleeping disorder. Improves the quality of sleep of secondary sleep disorders. |
| Li et al.13                   | Systematic review and meta-analysis | 205 patients between 29 and 41 years old | To determine the effectiveness of melatonin versus placebo in the treatment of sleep disorders. | 3 - 6 mg for 3 to 9 days. | Sleep onset latency, total sleep time and sleep efficiency. | Improves the quality of sleep of secondary sleep disorders.                                                                 |
| Malow et al.42                | Double-blind placebo-controlled clinical trial | 80 children and adolescents aged between 2 – 17.5 years old | To report the long-term effects of prolonged-release melatonin treatment appropriate for children on sleep, growth, body mass index and pubertal development. | 2, 5, 10 mg/day intake for up to 104 weeks, followed by a 2-week placebo period to reassess withdrawal effects. | Infant sleep was assessed at each visit during the one-year study period using the Compound Sleep Disorders Index, which scores the frequency and duration of the participant’s sleep habits in the previous month (six habits: sitting at bedtime, sleep induction, waking up at night, resettlement, time of wake up, and early sleep). | It is safe and effective for long-term treatment in children and adolescents with autism spectrum disorder and insomnia. There were no detrimental effects on children's pubertal growth and development and no abstinence or safety problems related to the use or discontinuation of the medication. |
| Low, Choo & Tam47             | Systematic review           | Patients aged 18 - 65 years old | To summarize all available systematic reviews and meta-analyses that investigate the effectiveness of melatonin and melatonin agonists in primary insomnia disorders. | 0.3 - 75 mg, 3 days to 6 months | Actigraphy, waking time during sleep, awakenings, day to night sleep ratio. | There was a statistically significant improvement in latency and total sleep time, with a lack of consensus on whether these are clinically significant. |

CRA-melatonin: melatonin of continuous release and absorption; DSWPD: Delayed Sleep-Wake Phase Disorder; ICU: Intensive Care Unit; IR-melatonin: melatonin of immediate release
of melatonin used in these studies were equal to or higher than the doses indicated for healthy elderly people, however, the doses were lower compared to those used or studied in populations without dementia. Several different mechanisms are likely to cause sleep disorders in patients suffering from dementia, some of which may be related to circadian misalignment. Achieving full melatonin’s chronobiotic effect in these circumstances can take several months. Therefore, it is possible that some patients respond after longer periods of treatment with melatonin.

Overall, no significant adverse effects were reported in most studies; however, Besag et al. reported that daytime drowsiness and headache are among the most frequent related side effects. Similarly, Maras et al. stated that after 52 weeks of use of melatonin the most frequent treatment-related adverse events observed were fatigue and mood swings. Additionally, Myers et al. in a randomized clinical study, reported that a patient had an increased serum level (within the toxic range) of valproate while using melatonin concomitantly. According to the authors, clinicians must ponder possible interactions of melatonin with antiepileptic drugs. Since melatonin is also metabolized by cytochrome P450 enzymes (CYP1A2, CYP1A1, and CYP2C19), the concomitant use of melatonin with antiepileptic and antidepressant drugs can potentially cause drug interaction. Consequently, the metabolism may be reduced leading to longer drug action time, which can cause severe sedation.

A formulation containing melatonin, vitamin B6, California poppy extract, passion fruit extract, and lemon balm extract was tested on patients of both genders between 20 and 75 years old who had moderate insomnia. There was a statistically significant improvement in sleep quality during the two-week treatment period with no serious adverse events being reported, suggesting that this combination of assets is beneficial for mild to moderate insomnia.

As for the doses, 0.1 to 0.5 mg is recommended for sleep-related rhythmic movement disorder, 1 to 5 mg for sleep disorders, and 3 to 10 mg for neurological diseases. These recommended doses should be taken daily, in a single dose at night, one hour before the usual bedtime. However, there is no established minimum or maximum effective dose for each use. Among the advantages of melatonin described in the included studies are its favorable safety profile, good toleration, and no addiction potential when administered for long periods. The adverse reactions commonly found in the reviewed studies included headache, nasopharyngitis, back pain, arthralgia, nausea, dizziness, and restlessness. Additionally, melatonin may have advantages attributed to its anxiolytic and hypnotic effects, and low toxicity levels, as reported by Madsen et al. while investigating the melatonin toxicity in patients with depressive, anxiety, and sleep disorder symptoms.

In Europe, the drug Circadin, which has a prolonged release formulation containing 2 mg of melatonin, has been marketed since 2008 as an innovative treatment for primary insomnia in patients aged 55 years and older who have sleep disorders characterized by poor quality of sleep. As a food supplement, it has not been evaluated or approved by the United States FDA to prevent or treat any diseases.

In Brazil, there is no registration of drugs with melatonin as the active ingredient, therefore, its sale is prohibited in drugstores and national websites. Although melatonin is used in some countries as an ingredient in food supplements, this substance is not authorized for use in food supplements in Brazil. Additionally, according to the rules of Ordinance SVS / MS no 344/1998, melatonin is not an asset subject to special control, however, its commercialization is restricted to compounding pharmacists which must acquire the product from the company Active Pharmaceutica, have operation authorization granted by the ANVISA, and follow the current guidelines of Good Handling Practices. Additionally, prescription from a legally qualified professional is required, and the prescription must contain the composition, the pharmaceutical form, the dosage, and directions for use.

In conclusion, it is evident from the identified studies that melatonin can be used in specific dosages according to age for sleep disorders, jet lag, insomnia in children with neurological disorders. Exogenous melatonin has emerged as an alternative therapy that can be used in sleep disorders. According to the evidence found, melatonin has not demonstrated toxicity or severe adverse effects, nor dependence even at high doses, demonstrating that its use is safe for treating young and elderly patients. However, despite the findings discussed, further investigations are needed in order to assess the dosages required for each age group, as well as dosages’ safety profile.

Currently, no melatonin drug has been approved for use by regulatory bodies or legislation in Brazil, and the available
information only determines the pharmaceutical form, excipient substance, and general guidelines for melatonin handling. Guidelines addressing the accurate use of melatonin to support clinicians and pharmacists in the treatment decision-making for sleep disorders is required. In addition, as melatonin is a relatively new drug, pharmacovigilance is essential, as it is up to health professionals to report any adverse events to the authorities.

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