OPTIMIZING STABILITY OF PLANT EXTRACTS USING LYOPHILIZATION

Abstract

The growing need for bioactive metabolites necessitates an efficient method of extracting and processing diverse plant extracts. The majority of the existing literature focused on determining the influence of extraction procedure on plant extracts. However, the process used to dry plant extracts after extraction is also critical. The "gold standard" drying plant extracts for is lyophilization, also known as freeze drying, which is crucial for maintaining their quality and increasing their shelf life. A literature search using Pub Med and Google Scholar was done, and a summary of all the particular issues associated with lyophilization was drawn up. Due to the low process temperature, freeze-drying is used as a microencapsulation technology and is especially ideal for the entrapment and protection of delicate bioactive chemicals. The zeta potential, poly dispersity index (PDI), and minimum particle size are typically used to optimize the stability of lyophilized extracts. In this review, the impact of lyophilization on the stability of plant extracts has been explored.

Keywords: Lyophilization, Stability, Plant extracts, Encapsulation, Zeta Potential, Poly dispersity Index.

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I. INTRODUCTION

Many drug discovery efforts have used plant extracts as a significant source of bioactive chemicals and some significant medications have been extracted and discovered from plants [1]. Small biomolecules, biologics produced by plants and a recently established third category of medications known as phytopharmaceuticals makeup this class of medications [2]. Around 80% of the world's population, according to the World Health Organization (WHO), relied on medicinal plants as their main source of treatment in 1985 [3]. Various medicinal preparations, including powders, tinctures, tablets, etc., use plant extracts in crude form or standardized fractions [4]. Plant extracts contain a variety of bioactive substances with strong biological effects, such as polyphenols, alkaloids, and terpenes [5]. An adequate strategy is needed to effectively extract and process different plant extracts due to the growing demand for these phytochemicals [6]. Polar solvents (such as water and alcohols), intermediate polar solvents (such as acetone and dichloromethane), and nonpolar solvents (such as n-hexane, ether, and chloroform) are frequently used in the extraction of medicinal plants. Optimal extraction and the ability to maintain the stability of the chemical structure of desired substances are two qualities that define a good solvent [7]. The image illustrate show these bioactive chemicals are now subjected to a variety of thermal, non thermal, and physicochemical variables, including temperature, pH, pressure, sonication, radiation, electric field, humidity, oxygen, and light, as aresult of advancements in extraction technology (Figure 1) [8]. These processing and storage parameters have been thoroughly investigated by researchers, who have found that they have a considerable impact on the concentration, bioactivity, functionality, bioavailability, and overall stability of the active chemicals [9].

Factors Affecting Extraction Technology									
Humidity	Solvent pH	Temperatur	e Extraction technique		Extraction time				
	PH	~	6 ⁶ 6 9 ₉ 9		الم الم الم				
Agitation	Solvents	Pressure	Plant material	Material to solvent ratio					
			Selling Selling	Contraction Contraction					

Figure 1: Factors Affecting Extraction Technology

II. BACKGROUND

Enhancing the quality of plants and enhancing the safety, quality, and consistency of the finished products need the adoption of optimized high-quality agricultural and biomass processing processes [10]. The primary active substances utilized in the creation of natural products, which are often offered in the form of powders or tablets inorder to ensure good stability and appropriate dosage, are dried plant extracts high in bioactive metabolites [11]. Inorder to manufacture these natural compounds, it is crucial to optimize the extraction protocol, which comprises thesolvent choice, solvent to biomass ratio, extraction time, and extraction method choice [12]. The majority of the material that was published compared different extraction methods utilized for plant extracts. But it's also crucial to consider the process for drying plant extracts after extraction [13]. Especially because current research indicates that the drying technique used to prepare plant biomass prior to extraction influences their bioactivity and chemical makeup [14]. Similarly, the method of post-extraction drying used has an impact on the chemical and biological characteristics of plant extracts [15]. A mind map enumerating several drying techniques is represented as a figure (Figure2). Lyophilization is one of the more modern and inventive post-extraction techniques. The "gold standard" for drying plant extracts is lyophilization, also known as freeze drying, which is crucial for maintaining their qualityand increasing their shelf life [16]. With the fore mentioned context in mind, this review attempts to provide insight into the various ways that lyophilization can be used to maintain the stability of plant extracts.

DRYING METHODS FIRST GENERATION SECOND GENERATION THIRD GENERATION FOURTH GENERATION TRAY MICRO-WAVE TUNNEL DRUM • FREEZE HIGH VACUUM TRUCK-TRAY SPRAY OSMOTIC · RADIO- ROTARY FLOW FREQUENCY CONVEYER Non-thermal drying of Novel drying of liquid Thermal drying of liquid Thermal drying of solid and solid material and semi-liquid materials solid and liquid materials materials

Figure 2: Methods of Drying

III.METHODOLOGY

A PubMed and Google Scholar literature search was conducted using the key terms "lyophilization", "freeze-drying", "plant post-extraction methods", "lyophilized plant extract", "Stability studies" and "preserving bioactive compounds" to gather the information for this manuscript. An outline of all the special issues exclusive to lyophilization was prepared. All of the data cited has been written entirely in English.

1. State of the Art: Lyophilization: Lyophilization in the pharmaceutical industry has undergone continuous growth and steady expansion aswe approach a new century. In the freeze-drying process, water is sublimated directly from the solid state (ice) to the vapor state, omitting the liquid state, and is subsequently reabsorbed from the "dry" layer [17]. Although lyophilizes are a more recent invention than freeze-drying, they have only been around for about 100 years [18]. The procedure preserves the dried product's quality (including its biological, dietary, and oil factory qualities) [19]. Lyophilization is always used to increase the stability of a product that is sensitive to moisture or to make the product simpler to store or transport [20]. Three basic processes are involved in the

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pharmaceutical lyophilization, as depicted in the picture (figure3). The first step is to freeze the product in solution to create a matrix of ice along with additional crystallizable excipients while concentrating other solutes and the active pharmaceutical ingredient (API) inside the interstitial voids [21]. Aqueous medicinal formulations can be frozen at temperatures below -35°C [22]. The second phase is primary drying, which involves submerging ice in a vacuum at a low temperature. Traditionally, this process is performed at chamber pressures of 40-400 Torr at shelf temperatures ranging from -30°C to -10°C [23]. After the primary freeze-drying process is completed and all ice has sublimed, bound moisture remains in the product. Although the product seems dry, the residual moisture level could be as high as 7-8% [24]. As a result, a last process called secondary drying at a warmer temperature is required to remove the residual moisture content to optimal levels. This is known as 'Isothermal Desorption' because the bound water is desorbed from the product [25]. By increasing the shelf temperature above ambient levels, this process is completed. The shelf temperature can be increased to 15-300°C to allow for vacuum-induced desorption of water molecules [26]. Four standard parts are typically included in freeze-dryers: a drying chamber, a vacuum pump, a heat source, and acondenser [27]. The tools and procedures are created to guarantee that the sterility of the product is preserved throughout the lyophilization process [28].

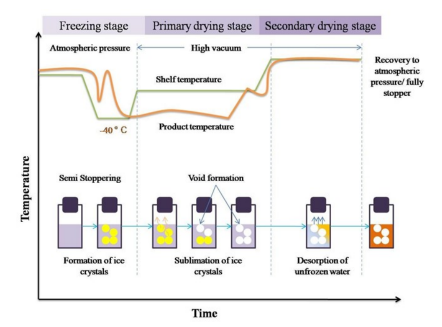


Figure 3: Lyophilization Principle and Process Lyophilization and its Multidisciplinary Application

Lyophilization is a common technique used to concentrate extracts or stabilize biological samples before storage [29]. On a laboratory and industrial scale, the freezedrying method is mostly employed for delicate, temperature-sensitive plant materials [15, 30]. By using the freeze-drying method, plant samples are highly effective in removing water while retaining bioactive components, such as antioxidants [31]. The freeze-drying procedure is frequently employed to preserve and stabilize biological materials because of its benefits [32]. It guards against both the spread of microbes and the deterioration of biological materials linked to the activity of bacteria [33]. In order to prepare plant and food samples for the identification of physiologically active chemicals, the freeze-drying method are also used [34]. It is used to preserve the chemical makeup of the plant material while stabilizing, enhancing, and/or extending its life [35]. Due to the low process temperature, freeze-drying is used as a micro encapsulation technology, making it especially ideal for the entrapment and protection of delicate bioactive chemicals [36]. In freeze-drying-based encapsulation, the target component and the encapsulating ingredients create an emulsion solution, which is then used to create microcapsules using the freeze-drying method [37]. Additionally, due to the few stages required, the freeze-drying technique is simpler than other micro encapsulation techniques [38].

2. Effect of Lyophilization on Stability of Plant Extracts: In order to maximize zeta potential and minimize polydispersity index (PDI), lyophilized extract stability istypically optimized [39]. The effects of lyophilization technique on stability of plant extracts are enlisted as table (Table1). The poly dispersity index is a size-based indicator of a sample's heterogeneity [40]. Size distribution in a sample, as well as aggregation or agglomeration of the sample during isolation or analysis, canal lead to poly dispersity [41]. Good homogeneity is indicated by a PDI value below 0.3 [42]. The zeta potential's size reveals the strength of electrostatic attraction between nearby, similarly charged particles in dispersion [43]. For sufficiently small molecules and particles, a high zeta potential will impart stability, which means that the solution or dispersion will fend off aggregation. Zeta potential readings exceeding 30 mV are often indicative of high stability [44]. Plant extracts' zeta potential is influenced by the extraction process and, consequently, by the extracted chemicals [45]. The thickness of frozen material and its surface area are additional factors. The rate of lyophilization increases with the frozen material's surface area. In contrast, the rate of lyophilization is slower the thicker the frozen substance. The ability of a sample to absorb and transfer heat to the surface undergoing sublimation is influenced by sample thickness [46].

S.No	Plant Extract	Stability studies	Source
1	Serpylliherba	Zeta potential at pH6:- 18.7±0.7 mV	[47]
2	Alliumcepa L.	Particle size:177.73 nmPDI:0.45	[48]
3	Artemisiaabsinthium	Zeta potential:-11.9 Particle size: 253.8 nmPDI:0.258	[49]
4	<i>Betavulgaris</i> L. Leaf hydro alcoholic extract nanogel	Zeta Potential:+28.8mV Particle size:247nm PDI:0.259	[50]
5	Anoectochillus burmannicus Ethanolic Extract-Synthesized Selenium Nanoparticles	Zeta Potential:-24.5±1.9m VPDI:0.366±0.074	[51]

6	<i>Hippophaes rhamnoides</i> aqueous extract-containing nanoemulsion	Zeta Potential (mV):- 30.11±2.02 Droplet Size (nm):183.07± 9.53 PDI:0.295± 0.045	[52]
7	Punica granatum L. Loaded nanophytosomes	Particle size:166.70– 144.40nm PDI:<0.5	[53]
8	Chitosan-Coated- <i>Aloeperryi</i> - Solid Lipid Nanoparticles(C- ALP-SLNs)	Zeta Potential: 13.6 ± 1.1 mV Particle size: 173.6 ± 11.3 nm PDI: 0.21 ± 0.02	[54]
9	<i>Eupatoriumadenophorum</i> Leaf extract-Silver Nanoparticles	Zeta Potential:- 33.4mV Particle size:117.75nm	[55]
10	<i>Calotropis gigantean</i> -loaded poliglusam-silver nanomatrices	Zeta Potential:42.42mV Particle size: 178.5 nmPDI:2.97	[56]

IV. CONCLUSION

Researchers have been inspired by the increased demand for plant extracts to create novel drying processes that preserve high concentrations of essential oils and other physiologically active substances without compromising the dried plant material's flavor, color, or texture. Lyophilization appears to be a useful approach for conserving the volatile profile of plants, although its effects depend on the species of plant (the location of volatile accumulation, the type of volatile components, etc.). The lyophilization process' technical specifications also have a big impact on the end product's quality. In order to more accurately examine the therapeutic benefits of plant extracts, additional systematic research is required on the effects of freeze-drying in comparison to alternative preparation methods, such as alcohol extractions of fresh plant material.

- **Competing Interests:** The authors declare no competing interests.
- Ethics Approval: There are no studies on human volunteers or animals in this article.
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