Influence of Particle Size and Particle Deposition of Inhaled Medication in Lung Disease: A Comprehensive Review

Kiran R. Dudhat1,2*, Harsha V. Patel3

1School of Pharmacy, R K University, Kasturbadham, Rajkot, Gujarat-360020, India.
2Department of Pharmaceutics, Gujarat Technological University, Chandkheda, Gujrat, India
3Indukaka Ipcowala College of Pharmacy, Vithal Udyognagar, Beyond G.I.D.C., New Vallabh Vidyangan, Gujarat - 388 121, India

ABSTRACT

Drug particles less than 5 µm have the greatest probability of deposition in the lung, whereas those less than 2 µm tend to be concentrated in the alveoli. A large proportion of particles within the 2-5 µm range are present in the dose released from the inhaled drug, providing a relatively even distribution across the lungs. The efficient need for inhaled therapy highly depends on the essence of the method of drug delivery and the patient’s ability to correctly use the system. A large range of inhaler products, each with positive and negative aspects, are on the market. It facilitates the administration of a lower dose; there is a quicker onset of action and less severe side effects. The deposition of the inhaled drug in the lung is dependent on particle size, inhalation technique and the type of inhaler device. Importance of particle size distribution and Particle aerodynamic diameter. Influence of environmental humidity on particle size Particle deposition in the airways, Methods to identify drug deposition in lungs. Physiological factors which affect the therapeutic efficacy of pulmonary delivery drugs. The nano and micro size particles is a mainstay of treatment for a variety of pulmonary diseases because they provide a platform to deliver drugs directly reliably and inexpensively to the disease site, thus allowing for a minimum amount of drug to be used and minimize side effects.

BACKGROUND

A non-invasive, rapid, and efficient approach to administering therapeutic agents locally and systemically is the pulmonary route of administration. On the other side, since the lungs have a large surface area available for absorption and ample vasculature, inhalation often has a great opportunity for systemic transmission.[1] The delivery of a drug to its target site might result in a fast onset of action, which is often desired. Compared to oral or parenteral administration, smaller dosages can be given locally, minimizing the risk of potent side effects and lowering treatment expenditures.[2,3] When a drug is poorly absorbed orally (e.g., sodium cromoglicate)[4] or rapidly metabolized orally (e.g. isoprenaline), the pulmonary route is also effective.[5] Although the lung has considerable metabolic potential, avoiding first-pass (presystemic) metabolism in the liver may be preferable.[2]

Compared to the liver and gastrointestinal tract, drug-metabolizing enzymes are smaller in the lungs. These characteristics generate conditions well suited to the successful absorption of drugs, providing a possible route for delivering systemic drugs.[6,7] Pulmonary drug delivery, however, is a difficult administration route. Next the efficacy of inhalation therapy depends on the site of the drug's deposition in the lung. Inhaled drug deposition is a complex process dependent on the lungs' anatomy and physiology, the drug's physicochemical characteristics, the design and characteristics of the formulation, and the type of delivery device used for administration.[8] The movement and deposition of particles in the lungs are highly affected by the airway geometry along the

*Corresponding Author: Kiran R. Dudhat
Address: School of Pharmacy, R K University, Kasturbadham, Rajkot, Gujarat-360020, India.
Email: kichupatel@gmail.com, kiran.dudhat@rku.ac.in

Relevant conflicts of interest/financial disclosures: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2022 Kiran R. Dudhat et al. This is an open access article distributed under the terms of the Creative Commons Attribution-NonCommercial-ShareAlike 4.0 International License which allows others to remix, tweak, and build upon the work non-commercially, as long as the author is credited and the new creations are licensed under the identical terms.
respiratory tract. In the alveolar zone, the main site of absorption, only particles of a particular size (usually 5 μm) and shape can deposit. 

**Anatomy and Physiology of Lungs**

The lung is the organ of outward breathing, where oxygen and carbon dioxide are exchanged between blood cells and inhaled air. The design of the aviation pathways also efficiently prevents the entrance of foreign airborne particles, including bacteria, and speeds up their evacuation.

It is possible to consider the respiratory tract as containing the conducting (central) regions (trachea, bronchi, bronchioles, terminal, and respiratory bronchioles) and the respiratory (peripheral regions (respiratory and alveolar bronchioles), although there is no definite demarcation between them (Fig. 1). The nose, throat, pharynx, and larynx form up the upper respiratory system, whereas the trachea, bronchi, bronchioles, and alveolar regions make up the lower respiratory tract. The airways will be specified by a symmetrical design in which each airway splits into two identical divisions or generations. The trachea (generation 0) splits into two main bronchi (generation 1). The right bronchus is larger and leaves the trachea at a lower angle than the left, making it more likely to accept inhaled particles. Further branching of the airways eventually led to the formation of terminal bronchioles. These are divided into respiratory bronchioles that interact with alveolar ducts that lead to alveolar sacs and alveolar sacs that communicate with respiratory bronchioles (generation 2, 3). These contain approximately $2 \times 10^8 - 6 \times 10^8$ alveoli, producing a surface area of 100 m$^2$ to 140 m$^2$ in an adult male.

The blood barrier between the alveolar space and the pulmonary capillaries is exceedingly thin to allow for fast gas exchange. The alveoli are mucus-extracted and have a much flatter epithelium, 0.1-0.5 μm thick, simple squamous form. A floor-active component that consists of phospholipids, called lung surfactant, is coated with the alveolar floor; its location is later stated. Conducting airways are lined in ciliated epithelial cells. Insoluble particles deposited on the walls of the airways in this area capture the mucus, which is then carried up from the lungs by the beating cilia to the throat and swallowed.

**Absorption of Particles in Respiratory Tract**

Small molecule medications and several therapeutic peptides and proteins are naturally permeable to the pulmonary membrane. The broad barrier to the absorption of inhaled drugs is the epithelium of the lung. In the trachea, it is dense (50–60 μm), but in the alveoli, its thickness decreases to 0.2 μm. When moving distally from the trachea, bronchi, and bronchioles to the alveoli, cell types and shape change is significant. Macromolecules are more sensitive to the lungs than any other bodily entrance site. A few peptides have demonstrated very high bioavailability through the pulmonary route, especially those chemically altered to inhibit peptidase enzymes. If they are strongly cationic, small molecules can show prolonged absorption. Whereas rapid molecular absorption in the lungs has a variety of medicinal applications, there are times when slowing the absorption rate of inhaled small molecules is necessary, either to keep them working locally in the lungs or to regulate their absorption throughout the body. Very insoluble Molecules that slowly dissolve from an inhaled particle might reside in the lungs for hours or even days.

The drug absorption rate and quantity differ over the length of the respiratory tract. For example, different areas of each region (about 2 m$^2$ conducting airways but about 140 m$^2$ alveolar surfaces) are influenced by absorption in different regions.

The airways and alveolar areas have variable epithelial thickness and cell populations. A mucus gel covers the airway epithelium, while a surfactant layer coats the alveolar surface. Drug clearance from the trachea and bronchi is mainly mediated by ciliated cells and mucus, while macrophages largely handle clearance from the deep lung. In conjunction with mucus, the ciliated cells form a significant route for drug clearance from the trachea and bronchi, while macrophages are critical for deep lung clearance.

The amount of drug deposited and diffused inside the lungs was determined by an aerosol’s overall therapeutic efficacy. These processes are a physical barrier to aerosolized drug delivery to the airways. A complete explanation of the role of each physiological zone about ultimate pharmaceutical absorption needs a solid understanding of lung anatomy and physiology.

Pneumocytes line the surface of the alveoli in two types: type I pneumocytes, which are thin squamous cells that form an important part of the capillary gas exchange barrier, and type II pneumocytes, which are bigger cuboidal cells that generate lung surfactant and are more diffuse than type I cells. Alveolar (phagocytic) macrophages scavenge and transport particulate matter to
the lymph or mucociliary escalator, accounting for around 3% of the alveolar region’s cells.\[^{30}\]

Larger particles (5-10 \(\mu\)m) are deposited in the oropharyngeal area and larynx because of impaction. Particles of a diameter of 1 to 5 \(\mu\)m are normally seen in the tracheobronchial tract. In the alveoli and narrow conducting airways, gravitational sedimentation deposits particles with a diameter of 0.1-1 \(\mu\)m. Particles smaller than 0.1 \(\mu\)m are not deposited and are ejected during exhalation (greater than 0.1 \(\mu\)m are deposited and expelled).\[^{33}\]

Fig. 2 shows particle size-dependent deposition in various parts of the respiratory tract. To investigate the science of bronchodiulator particle size effects, the research has already assessed regional airway drug deposition with the simultaneous calculation of clinical response. The system uses various particle sizes to transmit the target effect or cells to inhaled lung regions.

**Mechanisms of Drug Absorption from the Lungs**

The lung contains many of the absorption processes involved in other routes of administration that occur in organs. In general, paracellular or transcellular absorption of inhaled drugs may be possible. Paracellular absorption occurs via close junctions, claudine, and occludine integral proteins that stretch between lung epithelial cells in the paracellular space.\[^{32}\]

Research findings have also shown that apical to basal trans-epithelial electrical resistance, which shows the degree of cell tightness, decreases from the tracheal area to the distal airways until the alveolar region increases again. Therefore, paracellular absorption in the distal bronchioles is most likely to occur. Several hydrophilic drugs with relatively small molecular weights, such as insulin (Mol. Wt.: 5808 Da), have been documented to be absorbed into the lungs through paracellular transport.\[^{22}\]

A few other methods can improve the paracellular transport of drugs, for example by reversibly decreasing the tightness of paracellular junctions by administering compounds such as chitosan, allowing larger molecules to move across.\[^{33}\]

Transcellular transport, in which the medication must diffuse into the cells to be absorbed, forms a major part of drug absorption through the lungs.\[^{32}\]

Transcellular transport, in which the medication must diffuse into the cells to be absorbed, forms a major part of drug absorption through the lungs.\[^{32}\]

Transcellular transport frequently requires transporter molecules that are expressed on the cell membrane surface. In relation to intestine, liver, and kidney transporters, there is a scarcity of information about lung transporters.\[^{35}\]

Most of the other transporter express studies were done in vitro, which means that the degree of expression or distribution of transporters in vivo may not be accurately described. Furthermore, nothing is known about the extent to which such transporters play a role in the absorption kinetics of several drugs.

The solute carrier and ATP binding cassette transporters are two major transporters found in lung cells.\[^{36}\]

Via organic cation transporters and organic anion transporters, the solute carrier can transport organic cationic or anionic molecules. They can either enhance or impede the absorption of the drugs, depending on the location of expression of such receptors, either on the apical side at the lumen of the airway or on the basolateral side facing the endothelium of the blood capillaries. For such transporters, an immense diversity of substrates makes these receptors an important problem to consider during dosing calculations. Vesicular transport, which involves the creation of invaginations in the cellular plasma membrane that later separate into individual vesicles that engulf the particles within, is another potential absorption mechanism.\[^{37-39}\]

Depending on the particle size, vesicular transport can be mediated by either caveolin- or clathrin. Caveolin-mediated transport usually includes particles of less than 120 nm in size, while clathrins transport larger particles of 150-200 nm in size.\[^{37}\]

**Lung Surfactant**

Pulmonary surfactant, a lipoprotein complex composed of 90 percent lipid and 10 percent protein synthesized, secreted, and recycled by type II epithelial cells in the alveoli, lines the pulmonary airways. In minimizing surface tension and being host protection against inhaled pathogens and particles, the surfactant film of the lung plays a dual role.\[^{40,41}\]

Alveoli are stabilized against collapse by reducing the alveolar surface tension at the air-liquid interface, thereby maintaining a wide surface area for gas exchange.\[^{42}\]

The surfactant also enables oxygen penetration into the blood and through the lung surface lining. It would be extremely difficult to breathe without the lung surfactant, as oxygen diffusion through the lung surface lining would be hampered.\[^{43,44}\]

Lung surfactants also experience anti-inflammatory and antioxidant effects. In addition, pulmonary surfactants allow the movement of accumulated particles to the bronchial tree’s upper airways. However, interactions have been identified between the lung surfactant phospholipids and inhaled drugs.\[^{45}\]

Lung surfactant has been shown to increase
the solubility of steroidal drugs (glucocorticosteroids), which have affected their residence time in the lung, with anti-inflammatory and antioxidant impact. Other studies have shown that certain antibiotics can impair pulmonary surfactant function. These reactions between the antibiotic and the lung surfactant should also be closely examined before inhalation antibiotics are administered. Furthermore, potential interactions between deposited nanoparticles and lung surfactants can affect the role of biophysical surfactants, the metabolism of surfactants and the clearance of particles, or trigger toxicity caused by particles. It is reported that in the presence of a sufficient amount of aerosolized insoluble particles, the activity of the lung surfactant is decreased (e.g., polymer microparticles). This can disrupt the surfactant’s physiological function, including the retardation of particle clearance from burdened lungs. Lung surfactants can allow large molecules to accumulate, such as protein therapeutics, which may increase alveolar macrophages’ intake and digestion. They become enveloped by a lung surfactant monolayer as aerosol particles lodge in the lung. These cell - derived particles are quickly digested and then removed from the alveolar zone by macrophages. The certain new studies indicate that the diffusion of the medication out of the alveoli could be hindered by the lung surfactant. The addition of exogenous surfactant into the inhaled mixture expands the distribution further into the lung lumen of drug particles.

**Pulmonary Disease**

A wide variety of persistent pulmonary conditions have been implicated in lung diseases such as chronic obstructive pulmonary disease (COPD), cystic fibrosis, extreme progressive pulmonary hypertension, pulmonary tuberculosis, bacterial and fungal pulmonary infections, asthma, lung cancer, idiopathic pulmonary fibrosis (IPF), and multiple interstitial pulmonary diseases. Such diseases are therefore chronic and sometimes lethal; it takes longer to treat them. Any of them are not totally healed and no treatments have been found to be effective in completely recovering lung functions. The IPF that is commonly known is idiopathic pulmonary fibrosis, which makes up 45% of IPF patients. The frequency of IPF and COPD, irrespective of sensitivity to common environmental risk factors, rises with age. It is currently estimated that about 300 million and 210 million people worldwide suffer from common diseases such as asthma, IPF and COPD, respectively. Be that as it may, pulmonary tuberculosis remains the top irresistible executioner around the world, with 10 million individuals becoming sick with TB in 2018 causing over 3 million deaths. Conventional pharmacotherapy for chronic lung disorders can be broken into a few categories according to clinical agent types. A variety of antibodies, genetic compounds, peptides, and chemical drugs have been treated for persistent lung diseases (e.g., siRNA, miRNA, and shRNA). Drug Delivery to the Pulmonary System

Drug distribution to the lung system was performed by encapsulation of the drug to be administered in microparticles with a size range between 0.5 and 10 microns, ideally between 2 and 5 microns, formed by a drug-releasing substance with a pH greater than 6.4. In a desired embodiment, the drug distribution mechanism is based on the development of microparticles that are stable at a pH of 6.4 or less and unstable at pH of greater than 6.4, or which are stable at both acidic and basic pH, but which are unstable at pH between about 6.4 and 8. Many types of components can also be used, include biodegradable natural and synthetic polymers, such as proteins, polymers of mixed poly (hydroxy acids), alginate, and amino acids (proteinoids). The microparticles were changed in another embodiment only after reaching the targeted cells to effect targeting to particular cell types.

**Table 1: Targeted Formulations with size particles**

<table>
<thead>
<tr>
<th>Sr no</th>
<th>Drug</th>
<th>Disease</th>
<th>Formulation</th>
<th>Targeted site</th>
<th>Particle size</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Anticholinergic drug</td>
<td>COPD</td>
<td>Pressurized MDI</td>
<td>Pulmonary</td>
<td>0.8 to 5 μm</td>
<td>[64]</td>
</tr>
<tr>
<td>2</td>
<td>Ciprofloxacin</td>
<td>cystic fibrosis and COPD</td>
<td>DPI</td>
<td>Pulmonary</td>
<td>3.2 to 3.4 μm</td>
<td>[65]</td>
</tr>
<tr>
<td>3</td>
<td>Paclitaxel</td>
<td>lung cancer</td>
<td>DPI</td>
<td>Pulmonary</td>
<td>2 to 4 μm</td>
<td>[66]</td>
</tr>
<tr>
<td>4</td>
<td>Sodium cromoglycate</td>
<td>Asthma or COPD</td>
<td>DPI</td>
<td>Pulmonary</td>
<td>&lt;5 μm</td>
<td>[4]</td>
</tr>
<tr>
<td>5</td>
<td>Dehydroepian-drostone</td>
<td>Asthma or COPD</td>
<td>Nebulizer</td>
<td>Pulmonary</td>
<td>&lt;5 μm</td>
<td>[67]</td>
</tr>
<tr>
<td>6</td>
<td>Tilorone and Pirfenidone</td>
<td>IPF</td>
<td>Ultrasonic nebulizer</td>
<td>Pulmonary</td>
<td>&lt;5 μm</td>
<td>[68]</td>
</tr>
<tr>
<td>7</td>
<td>Paclitaxel</td>
<td>lung cancer, COPD</td>
<td>Nebulization</td>
<td>Pulmonary</td>
<td>4 to 6 nm</td>
<td>[69]</td>
</tr>
<tr>
<td>8</td>
<td>Tobramycin</td>
<td>Cystic fibrosis</td>
<td>DPI</td>
<td>Pulmonary</td>
<td>&lt;5 μm</td>
<td>[70]</td>
</tr>
<tr>
<td>9</td>
<td>Tacrolimus</td>
<td>Lung transplant rejection</td>
<td>DPI</td>
<td>Pulmonary</td>
<td>140 nm</td>
<td>[71]</td>
</tr>
<tr>
<td>10</td>
<td>Budesonide</td>
<td>Local anti-inflammatory</td>
<td>DPI</td>
<td>Pulmonary</td>
<td>1 to 5 nm</td>
<td>[72]</td>
</tr>
<tr>
<td>11</td>
<td>Ciprofloxacin hydrochloride and gatifloxacin hydrochloride</td>
<td>Respiratory infections</td>
<td>DPI</td>
<td>Pulmonary</td>
<td>&lt;5 μm</td>
<td>[73]</td>
</tr>
</tbody>
</table>
Influence of Particle Size and Particle Deposition of Inhaled Drug in Lung Disease

**Pharmacokinetics of Inhaled Drugs**

The inhaled mode of administration generates high local concentrations in the lungs and relatively low levels of systemic absorption when dosages with a high therapeutic ratio are delivered. The pharmacokinetic testing of drug absorption from the lungs offers a reliable and reproducible method of evaluating the various delivery mechanisms of inhalers, and some determining the bioequivalence of generic drug formulations. To determine the effects of the inhalation technique on drug delivery in vivo, measures of drug absorption from the lungs may also be applied. For example with salbutamol administered through a large quantity spacer, lung bioavailability is shown to have changed by factors such as the number of actuated puffs, inhalation-actuation delay and washing procedure. Differences between dry powder reservoir and pressurised metered-dose aerosol systems in drug delivery to the lungs directly translate into commensurate differences in clinical effectiveness for both inhaled corticosteroids and b2-adrenoceptor agonists for delivery. Pharmacokinetic assessment appears to have a defined purpose in the measurement of drug delivery to the lungs, and it provides useful information that is comparable to other methods such as radiolabelled deposition. Research into the pharmacokinetics of proven and novel drugs and delivery mechanisms over the next decade is anticipated with keen interest and, ideally, will provide a clearer understanding of how to improve the benefit-risk ratio for inhaled drugs.

**Distribution of Particulate Size and Aerodynamic Diameter**

The physicochemical characteristics of the drug like the formulation, the delivery/releasing method, and the patient are the four primary factors that influence drug particle deposition in the airways. The use of inhaled therapy has a few advantages over systemic (oral or intravenous) administration. As in the targeted inhaled dose (Fig. 3), the drug dose is requiring less as compared to conventional oral dose. Targeted dose with smaller particle size has improved and more drug effect with less side effect. Systemic larger dose is travel less to lung and give larger side effect. As in targeted inhaled smaller dose is goes into lung in higher amount and less in systemic circulation. Particles will distribute in lung as per its size and density. It facilitates the administration of a lower dose; there is a quicker onset of action and less systemic side effects. The deposition in the lung of the

---

**Table 2: Different In-Vivo Study of Pirfenidone with systemic and targeted dose**

<table>
<thead>
<tr>
<th>Subjects</th>
<th>Systemic dose (Oral)</th>
<th>Targeted dose (Pulmonary)</th>
<th>Targeted Route</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male C57B1/6 Mice (6-8 Week)</td>
<td>1197 mg/day</td>
<td>Efficacy Dose 10µg/kg</td>
<td>Intra-tracheal</td>
<td>[83]</td>
</tr>
<tr>
<td>Male Sprague Dawley Rates</td>
<td>200 mg/(kg.day)</td>
<td>20 mg/(kg.day)</td>
<td>Inhalation</td>
<td>[84]</td>
</tr>
<tr>
<td>Male Wistar Rates</td>
<td>25-800 mg/(kg.day)</td>
<td>50 mg/(kg.day)</td>
<td>Endo-tracheally</td>
<td>[85]</td>
</tr>
<tr>
<td>Male Sprague Dawley Rates</td>
<td>160 mg/kg</td>
<td>300 µg/rate 30-1000 µg/kg</td>
<td>Intra-tracheal</td>
<td>[86]</td>
</tr>
<tr>
<td>Normal healthy volunteers, Smokers (higher risk for intolerance), and Patients with IPF</td>
<td>801 mg</td>
<td>25, 50, 100 mg (nebulizer dose)</td>
<td>Inhalation</td>
<td>[87]</td>
</tr>
</tbody>
</table>

**Table 3: Marketed drug dose Differences of Systemic Dose and Targeted Dose**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Oral (Systemic) Dose</th>
<th>Inhalable (Targeted) Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Salbutamol</td>
<td>4 mg/day</td>
<td>100 µg/day</td>
</tr>
<tr>
<td>Bedomethasone diprop.</td>
<td>8 mg/day</td>
<td>250 µg BD</td>
</tr>
<tr>
<td>Fluticasone Propionate</td>
<td>5 mg QD</td>
<td>5 µg QD</td>
</tr>
<tr>
<td>Voriconazole</td>
<td>400 mg BD</td>
<td>40 mg BD</td>
</tr>
<tr>
<td>Terbutaline Sulfate</td>
<td>15 mg / day</td>
<td>0.5 mg / dose</td>
</tr>
<tr>
<td>Dapsone</td>
<td>100 mg BD</td>
<td>50-500 µg/day</td>
</tr>
<tr>
<td>Tacrolimus</td>
<td>2-4 mg/day</td>
<td>200 µg/dose</td>
</tr>
<tr>
<td>Sildenafil</td>
<td>60-300 mg/day</td>
<td>5 µg / day</td>
</tr>
<tr>
<td>Itraconazole</td>
<td>200 mg BD</td>
<td>10 mg, 20 mg/day</td>
</tr>
<tr>
<td>Ethionamide</td>
<td>250 mg /day</td>
<td>2 µg/day</td>
</tr>
</tbody>
</table>

BD- Two times in a day, QD- Four times in a day
inhalation. The technique of inhalation, and the form of inhaler device. The patient’s inhaler procedure using the system is an integral part of the deposition of the drug inside the lung. For the most efficient delivery of inhaled medications, the right inhaler technique is crucial.

The most essential physical feature of an aerosol for inhalation is its particle size. The particle size of aerosol measurement is carried out by the aerodynamic diameter of the aerosol (da), which is the physical diameter of an airborne unit of density with a velocity equal to that of the particle in consideration. It is used to standardize the particle size of an aerosol.\(^{88,89}\) When the scale is log-normally distributed, the geometric standard deviation (\(\sigma_g\)) is used to indicate the size distribution. For approximately aerodynamic diameter (da) for spherical particles,

\[
d_a = d_p \sqrt{p/p_0}
\]

Where \(d_p\) is the physical diameter, \(p\) is the particle density and \(p_0\) is unit density (i.e., 1 g cm\(^{-3}\)). When \(d_p\) is the mass median diameter, da is termed the mass median aerodynamic diameter.\(^{90}\) Porous particles are effectively transported to and accumulated in the lungs with large physical diameters on the order of 20 \(\mu\)m. Because of the porous or hollow nature of their structure, their low density means such particles have a limited aerodynamic diameter and are therefore brought deep into the lungs in the inspired air. Furthermore, large particles are less susceptible to aggregation than smaller ones, providing advantages of composition, and the particles are too heavy to be removed by alveolar macrophages from the airways.\(^{91,92}\)

**Influence of Environmental Humidity on Particle Size**

As a particle enters the respiratory system, the shift in relative humidity from baseline to high (99 %) causes condensation of water on the particle surface, which continues until the vapour pressure of water reaches that of the surrounding environment.\(^{93-96}\)

**Particle Deposition in the Airways**

The delivery by inhalation of therapeutic compounds provides a promising and efficient alternative to existing invasive techniques. The size and density of inhaled particles in various regions of the respiratory tract are primary factors in determining their efficiency of deposition.\(^{97}\) The application of an inhaled therapy agent within the lung has a significant impact on the efficacy of that treatment. In addition, it is important to distinguish the impact of the delivery mechanism from the pharmacological activity for precise evaluation during the drug development process.\(^{98}\)

The effectiveness of a therapeutic aerosol depends on its ability to penetrate and be retained in the respiratory tract. Aerosols need a size smaller than approximately 5 \(\mu\)m or 6 \(\mu\)m to penetrate to the peripheral (respiratory) areas, with less than 2 \(\mu\)m being preferable for alveolar deposition.\(^{86,99}\) In addition to the environmental changes in the previously mentioned size and the hetero disperse nature of inhalation aerosol size distributions, literature values for ‘respirable’ size differ and must be considered. In the upper respiratory tract, larger particles or droplets are deposited and are quickly removed by the mucociliary clearance process from the lung.\(^{100,101}\) As a result, systemic absorption of the drug becomes available and can potentially cause adverse effects. A sufficiently large corticosteroid aerosol can be accumulated in the mouth and throat, risk causing adverse effects, including oral candidiasis. The size of the aerosolized drug may be particularly significant in treating specific disorders where penetration to the peripheral airways is desirable, such as the treatment and prevention of Pneumocystis carinii pneumonia alveolar infection.

For particulate deposition in the lung, three mechanisms are mainly responsible: impact, gravitational sedimentation, and diffusion (Fig. 4(a) and 4(b)).\(^{102,103}\)

**Inertial impaction**

The airstream changes direction in the mouth, where a bifurcation happens in the respiratory tract.\(^{104}\) Instead of following the shifting airstream, particles within the airstream, having sufficiently high momentum, can impact on the airway walls. For large particles with a diameter greater than 5 \(\mu\)m and especially greater than 10 \(\mu\)m, the major method for deposition in the nose, mouth, throat, larynx, and major conducting airways is this mechanism particularly significant and prevalent in the upper airways.

---

*Fig. 4(a): Mechanisms for Particles Deposition in Lung*
Influence of Particle Size and Particle Deposition of Inhaled Drug in Lung Disease

The airflow decreases as the conducting airways are further divided, and the impact becomes a less prominent deposition process.\(^{[103,106]}\) The probability of impaction is proportional to

\[
\frac{V_t \cdot V \cdot \sin \theta}{\rho g r}
\]

...(2)

Where \(\theta\) is the change in the direction of the airways, \(r\) is the airway's radius, \(V\) is the airflow velocity, and \(V_t\) is the terminal settling velocity.\(^{[106]}\)

**Gravitational Sedimentation**

Gravitational sedimentation refers to particle settlement under gravity action, which occurs predominantly in narrow airways and alveolar cavities where there is a small distance to be filled by the particles before reaching the walls. A particle settling under gravity will achieve a constant terminal settling velocity \((V_t)\) from Stokes' law:\(^{[107]}\)

\[
V_t = \frac{\rho g d^2}{18 \eta}
\]

...(3)

Where \(\rho\) is the particle density, \(g\) is the gravitational constant, \(d\) is the particle diameter, and \(\eta\) is the air viscosity. As a result, the size and density of an inhaled particle and its residence duration in the airways affect gravitational sedimentation. Sedimentation is an important deposition mechanism for particles in size range from 0.5 µm to 3 µm, in the small airways and alveoli, for particles that have escaped deposition by impaction.\(^{[6,5,67]}\) Deposition by sedimentation in the airways and alveoli increases with increasing particle size and increasing particle residence time. Larger particles are primarily deposited by inertial impact and smaller particles by Brownian diffusion mechanism.\(^{[101]}\)

In the lower bronchial airways and the alveolar zone, where airflow is slower, particle deposition via sedimentation occurs. Particles with a diameter of 0.5-5 µm were that avoided impact in the upper airways and are deposited in the lower tracheobronchial and alveolar areas by deposition and impaction. Deposition in the tracheobronchial area is significantly more likely if the aerosol particle size is between 3 and 5 µm.\(^{[64,86]}\) If the particles are smaller than 3 µm in size, substantial deposition in the alveolar region could be expected. Particle sedimentation is governed by the (higher) gravitational force acting on the particles than the (lower) dragging force exerted by the airflow.\(^{[85,102]}\) With an increase in particle size and a reduction in flow rate, the rate of sedimentation deposition increases. For particles of a size greater than 0.5 µm, this mechanism is particularly important. The probability of deposition in cylindrical airways by sedimentation \((P(S))\) is calculated as:\(^{[102]}\)

\[
P(S) = 1 - e^{-4gC \rho d^2 L \cos \theta / 9\pi \mu R v}
\]

...(4)

Where \(g\) is the acceleration due to gravity, \(C\) is the Cunningham slip angle correction factor, \(\theta\) is the angle relative to gravity, \(\rho\) is the density of the particle, \(L\) is the tube length, \(d\) is the radius of the particle, \(\mu\) is the viscosity of fluid, and \(R\) is the radius of the airways.

**Brownian Diffusion**

Diffusion is the main mechanism of deposition caused by Brownian motion for particles less than 0.5 µm in size. With decreasing particle size and airflow rate, this motion increases and thus becomes an essential mechanism in the lower airways and alveolar region for particle deposition. Here, particles travel through the streamline from high to low concentration and deposit through interaction with the airway wall. The geometric rather than aerodynamic scale of the particles governs this process.\(^{[102,104,106]}\) When they collide with air molecules, nanoparticles deposit through diffusion due to the displacement. The probability of deposition in the cylindrical airways by diffusion \((P(D))\) is calculated as:\(^{[78]}\)

\[
P(D) = \sqrt{2KTC / 3\pi \eta d / R}
\]

...(5)

\[
Dif = \frac{kT}{3\pi \eta d}
\]

...(6)

Where \(k\) is the Boltzmann constant, \(R\) is the gas constant, \(T\) is the absolute temperature, \(\eta\) is the gas viscosity, \(\mu\) is the absolute temperature, \(\eta\) is the gas viscosity, \(\gamma\) is the gravitational constant, and \(d\) is the particle diameter.\(^{[109]}\) There are several barriers to optimizing pulmonary drug delivery system. The physical features of a particle, including size and density, together with the airflow in the lung, decide its final place of deposition. The airflow varies both spatially and temporally and is determined by the geometry of the lung, the mechanical properties (compliance and...
resistance) of the lung tissue, and the driving pressures produced by the diaphragm and intercostal muscles within the lung. The problem lies in the fact that it is very hard to test these effects in vivo, and that these factors will change all lung diseases will change these factors. In addition, it is difficult to determine the distribution of particle deposition in a non-invasive way. Subsequently, the deposition mechanisms' causes and consequences are covered under real conditions. This has contributed to a restricted understanding of different parameters' impact in clinical conditions on the deposition and distribution of particles.

Nuclear medical imaging has been the most commonly used experimental method for measuring regional particle deposition. A radionuclide contrast agent is inhaled into the lung, and the radiation released from such a agent is imaged to enable its local concentration to be measured. This allows for regional deposition measurement; however, spatial resolution is insufficient to associate deposition with precise airway positions. And thus, there is limited scope for investigating the fundamental drug deposition mechanisms using this process. A lack of experimental evidence has facilitated an emphasis on deposition computer simulation. Although this has proven to be effective in many instances, to accurately simulate particle deposition, both in healthy and diseased states, a thorough knowledge of airway geometry and pressure/flow inputs is needed.

**Methods to Identify Drug Deposition in Lungs**

It is necessary to calculate any variance in pulmonary deposition because the accurate calculation of the quantity of the drug to enter the lung makes it possible to check the dosage and the ability to link the dose to clinical effect. Responsive and accurate methods are needed to analyze therapeutic strategies in depth. A range of proven non-imaging and imaging methods may examine inhaled drug deposition.

**Non-imaging or Pharmacokinetic Methods**

The approaches that do not require imaging include the pharmacokinetic methods of charcoal-block and urinary excretion. Both pharmacokinetics (PK) measurement techniques will allow the relative pulmonary bioavailability of an inhaled medication to be quantified. The method of charcoal blocking uses the ingestion of charcoal to obstruct any systemic absorption through the gastrointestinal route, thus quantifying pulmonary bioavailability. By comparing 'area under the curve' data across various regimens, the urinary excretion method estimates systemic transmission, so it is useful to measure relative pulmonary bioavailability. Several research groups have used both PK data and lung imaging to measure deposition, and PK methods provide similar results to lung imaging data for total lung deposition. PK approaches have the advantage of not having a radiation dose over lung imaging methods, so it would seem like there is little need to go to an imaging system to test lung deposition. The key drawbacks of the PK data are that these methods may not be appropriate for all medicinal products and the lack of knowledge these methods provide about the regional deposition of the inhaled medicinal product inside the lung in vivo may be relevant concerning the clinical effect. Any drug formulation that can be radiolabelled adequately can be carried out through imaging studies. A recent workshop on the role of PK methods in establishing bioequivalence for inhaled drugs has reinforced the existing consensus opinion that the lack of reliable information on regional inhaled drug deposition is the limitation of the PK methods.

**Imaging Methods**

**Two-dimensional Gamma Scintigraphy**

The 2D planar approach has been employed in most research using scintigraphy to assess drug deposition. The lungs are photographed using one or more, gamma camera detector heads after inhalation of a radiolabelled aerosol to obtain a static view of the pattern of inhaled deposition. The standard procedure is for the study volunteer to stand, sit, or lie between two static gamma camera heads for an acceptable amount of time to obtain a simultaneous anterior and posterior image. The anterior and posterior images are then merged to generate a geometric median image. That image must then be corrected for background activity, radioactive decay, and imaging effects through various tissue densities, also known as tissue attenuation correction.

**Single Photon Emission Computed Tomography (SPECT)**

Except that the gamma camera system is set up to rotate around the patient/volunteer, gathering 360° data, SPECT uses similar concepts to 2D, planar imagery. Increased details on regional deposition and clearance are the main benefit of SPECT. The amount of time needed to obtain a SPECT study is longer than for 2D planar scintigraphy with acquisition times of 10-20 min, and when using a radiolabel that is easily cleared from the lung, this can trigger problems. In SPECT studies such as 99mTc-DTPA or 99mTc-colloid, slowly cleared radiolabels are favored for this purpose. Application of SPECT has to clinical trials in drug deposition, ventilation and perfusion defect evaluation, and other applications such as epithelial permeability analysis.

**High-Resolution Computed Tomography (HRCT)**

For the diagnosis and longitudinal evaluation of lung disease, data obtained from CT is very useful, but there has also been a rise in interest in using such data to understand inhaled drug deposition better. Detailed central airway anatomy, regional ventilation, and texture...
assessment of the lung, in health and in the presence of lung disease have been studied using HRCT data. To measure disease incidence and therapeutic action, CT density is also used. HRCT data can be used to produce central airway models for the mathematical modeling of aerosol deposition and has been used to demonstrate regional deposition, including deposition inequalities between left and right lungs, and to simulate broad edge points. The significance of human anatomical variation, even in the healthy population, and the impact on aerosol deposition are increasingly recognized. It can be seen that the inclusion of HRCT data is very helpful, but it has the apparent disadvantage of having a higher radiation dose than that used for SPECT imaging. For this purpose, a changed protocol is used conservatively and frequently to keep the radiation exposure to a minimum.

**Positron emission Tomography (PET)**

PET requires various scanners and uses a radiolabelling method that is distinct from that used in SPECT or 2D planar gamma scintigraphy. PET is often paired with CT (PET/CT) in modern scanners. Computer analysis will then create three-dimensional representations of a radiolabel-emitting positron. Some unique examples for inhaled drugs such as $^{125}$Insulin and $^{125}$Insulin, PET integrates the radiolabel into the formulation being tested and uses positron-emitters such as $^{11}$C, $^{18}$F. Although the resolution is higher than SPECT (4-6 mm), the images created are identical to those made using SPECT. However, the use of PET for inhaled deposition studies has some important logistical problems, most notably the cost, the complicated methods of radiolabelling, and the need to have a cyclotron on site within a short distance from the research centre. PET-CT enables the evaluation of high spatial resolution aerosol deposition, 3D ventilation, and perfusion data, and provides new insight into cellular inflammatory cell activity. However, the high cost, the difficulty of image analysis and the relatively few centres that can solve the technical problems surrounding using positron emitters as radio labels are opposed to PET.

**Factors Influencing the Therapeutic Efficacy of Drugs Delivered by the Pulmonary Route**

As shown in Fig. 5, many factors influence respiratory deposition and others explained below.

**Airway Geometry**

The lung deposition of aerosol droplets/particles is strongly affected by the airway design in the respiratory tree. Every bifurcation, branching and decrease in the lumen diameter of the airways in the respiratory tract encourages the probability of particle deposition by impact and reduces the therapeutic effect of the fraction of particles available. The structure of the pharynx and larynx affect the airflow in the trachea and bronchi. At bifurcations in the upper respiratory zone, the sudden decrease in the downwards diameter contributes to the generation of turbulent airflow that increases the deposition of particles in the upper airways.

**Inhalation Mode**

The inhalation mode greatly affects the degree and area of the respiratory structure's particle deposition. In the peripheral alveolar area of the lung, nose breathing increases the probability of deposition of fine particles (about 10 μm) since larger particles are preserved in the nose and pharynx. In comparison, mouth breathing increases the chances of deposition of coarse particles (about 10 μm) with in the upper tracheobronchial region. The time between inspiration and exhalation is increased by holding the breath, which encourages particle sedimentation at the periphery of the lung.

**Airflow Rate**

The regional deposition of aerosol droplets/Particles in the respiratory tract is significantly influenced by the change in the inspiratory airflow rate. By improving aerosol deposition in the oropharynx region and upper airways, rapid and turbulent airflow decreases the residence time of the particles in the airways, while slow inhalation contributes to deposition in the lower peripheral airways. Furthermore, the lower deposition proportion of fine particles is followed by an increase in the airflow rate and vice versa. At a very slow airflow rate, the inhalation of an aerosol reduces the probability of particle/droplet effects, reducing aerosol particles deposition in the upper respiratory tract and by sedimentation and diffusion.

![Fig. 5: Factors Influencing Respiratory Deposition](image-url)
targets the lower airways.\textsuperscript{[122]} Finally, increasing the number of tidal particles (the volume of air displaced between natural inspiration and expiration when there is no extra effort) increases the accumulation of aerosol particles in the lower bronchial and alveolar regions. These are all the key reasons why patients are advised to breathe slowly and deeply and hold their breath when inhaling a substance.\textsuperscript{[123]}

### Mechanism of Particle Clearance

The particles are either eliminated from the lungs after inhalation of aerosol particles through the lungs, absorbed into the blood / lymphatic circulation or degraded by drug metabolism.\textsuperscript{[124]} The various clearance mechanisms used to remove foreign particles in different regions of the respiratory tract are reviewed under the parts.

#### Muco-ciliary Clearance (MCC)

MCC represents an essential protective mechanism for eliminating the respiratory tract of insoluble inhaled particles and serves as a possible physical obstacle for drug penetration. In healthy subjects, most of the accumulated particles in the trachea-bronchial region of the respiratory tract are removed within 24 hours of inhalation. In contrast to the lower airways, MCC is predominant in the upper airways.\textsuperscript{[11,25,126]}

#### Mechanical Clearance

The removal of inhaled particles from the upper airways is aided by nose blowing, sneezing, coughing, and swallowing. This process occurs in the wider airways immediately after the deposition of particles. Coughing is automatically caused when a particle about $\leq 10 \mu\text{m}$ in size is inhaled. A high airflow rate is required for efficient cough clearance, and as this is only available in the upper airways, it is only in this area that it is successful. The cough becomes the main clearance method in respiratory illness conditions such as bronchitis, asthma or pneumonia where MCC becomes affected. Therefore, for the optimal drug effect, it is necessary to maintain aerosols at sizes of $\leq 10 \mu\text{m}$.\textsuperscript{[127,128]}

#### Enzymatic Degradation

Many inhaled drugs are substrates for the CYP450 enzymes found in the lung epithelia, despite the amount of degrading enzymes in the lungs being much less than that in the liver.\textsuperscript{[129]} Certain isoforms such as CYP2S and CYP2F have been recognized as lung-specific.\textsuperscript{[130]} Moreover, metabolic enzymes such as esterases and peptidases from Phase II are also expressed in the lungs. The concentrations of these enzymes vary greatly between the various types of cells lining the lung regions.\textsuperscript{[131]}

#### Alveolar Macrophages

The effectiveness of inhaled treatment can be severely restricted by the housekeeping role of alveolar macrophages.\textsuperscript{[10,132]} If another inhaled drug substance has low solubility and the particles stay in the alveoli for sufficient time, macrophages may be washed, decreasing the amount of medicinal product available for a therapeutic effect. The key obstacle to achieving controlled drug release is in the alveoli remains clearance by alveolar macrophages. Most of the substances used only to prepare particles are rigid and have all the physicochemical characteristics that make them an ideal target for macrophage absorption that can sustain the release of a drug for the long term.\textsuperscript{[133,134]}

### Lung Receptors

Most of the inhaled medications interact with pulmonary cell-expressed unique receptors. Pulmonary delivery performance can be increased by targeting individual cells with a low risk of systemic side effects. Therefore, understanding the numerous cellular receptors in the lungs provides the potential for pulmonary therapy to be more successful. The $\beta$-adrenergic receptors, muscarinic receptors, histamine receptors (H1 and H2), glucocorticoid receptors, leukotriene 1 receptors and prostacycline receptors are the most important receptor groups, none of which are uniformly distributed in the lungs.\textsuperscript{[111]} Most $\beta$-adrenergic receptors, certain bronchi, and the terminal bronchioles, are found in the epithelium of the alveolar walls. $\beta$-Adrenergic receptor agonists are medicines that function on the $\beta$2-adrenergic receptor, inducing smooth muscle relaxation and dilation of bronchial passages (salbutamol (albuterol), terbutaline and isoprenaline).\textsuperscript{[135]} In the submucosal glands and lung lymph nodes, the smooth muscles of the airways, bronchi, and alveolar regions have a higher density of M3 receptors, although they make up a lesser fraction. Methacholine acts to contract the smooth muscles through M3 receptors.\textsuperscript{[136]} The H1 and H3 receptors are both predominantly located in the human respiratory tract's bronchial smooth muscle. These receptors contribute to the mediation of increased vascular permeability and smooth muscle contraction in the respiratory tract.\textsuperscript{[137]} Many novel receptors have now been identified as potential targets for developing novel therapies for lung disease, including orphan receptors.\textsuperscript{[138]}

### Disease States

Bronchial obstruction and narrowing of airways occur in respiratory disorders due to the accumulation of mucus and inflammation. Cystic fibrosis is a genetic condition in which dense mucus is formed in large amounts by the epithelial cells of the lungs, decreasing the lumen diameter in all airways.\textsuperscript{[71]} Chronic bronchitis is associated with severe mucus generation, thickening of the alveolar wall, and small bronchi occlusion.\textsuperscript{[172]} Asthma is a chronic inflammatory condition due to constriction of the bronchial airways in response to a stimulus marked by airflow obstruction (pollutants, allergens, or exercise). A thickened mucus layer and subepithelial fibrosis can also result in this constriction in turn.\textsuperscript{[6]} These all-disease situations modify the geometry of the airways, resulting in...
variable airflow velocities, air resistance, and turbulence that affect the pattern of aerosol deposition in the lungs. The aerosolized drug is deposited more by the inertial impaction process in the upper airways rather than by a uniform distribution in the lungs. Particles are larger than 5 μm are primarily trapped and unable to enter the lungs in the oropharyngeal region, while particles smaller than 1-μm are generally exhaled without deposition. This altered deposition pattern may lead to drug efficacy loss. [70] Compared to healthy lungs, a significant rise in the deposition of ultrafine particles has been documented in the lungs of patients with bronchitis and asthma. Lung inflammation, oxidative stress and genotoxicity were found to cause inhaled ultrafine particles. [73] The mucociliary clearance (MCC) may be compromised by any accumulation of thick mucus in the airways, resulting in the patient becoming more vulnerable to infections of the airways and the latter may be required to change the absorption of drugs. In response to smoking, a higher rate of inhaled drug degradation has been reported, increasing the expression of metabolic enzymes. The degree of expression of either absorption or efflux transporters may be impaired by other diseases, thereby changing the bioavailability of inhaled drugs.

Effect of Age and Gender Difference

Because of the anatomical changes that occur at progressive ages that cause dissimilarities in airway geometry, the subject’s age affects aerosol deposition in the human lung. Research findings have also shown that, relative to adults, children have increased upper airway deposition of coarse particles, but that the overall volume of deposition is very comparable. [139,140] Compared to infants, healthy adults can have a greater amount of aerosol accumulated in the alveolar region due to their greater lung volume. [141]

Anatomical variations between males and females in the larynx and airways are attributed to gender discrepancies in aerosol deposition patterns, relative to males, with females having more upper airway deposition. Studies have shown that females have higher coarse particle deposition (>5 μm) compared to males at a comparable flow rate, while fine particle aerosols (0.5 to 3 μm) have similar deposition patterns irrespective of gender. [142,143]

RECENT TECHNOLOGIES OF PULMONARY DRUG DELIVERY

Dry Powder Inhalation for Pulmonary Delivery

The use of dry powder inhalation therapy to treat respiratory and systemic disorders has been improved by advances in pulmonary drug delivery technologies. The effectiveness of dry powder inhalation therapy depends on at least four variables: the physicochemical properties of the formulation materials, the configuration of the device, the dispersion mechanism of the powder and the manoeuvre of the patient’s inhalation. [123,144] Powder particles delivered by DPIs should have an aerodynamic diameter of between 1 and 5 μm in order to pass across the respiratory system and most possibly enter the lungs. [145] The aerodynamic diameter (daer) determines how deep the aerosol particles can deposit inside the airways. The aerodynamic diameter is defined as the diameter of a sphere of unit density with an equal terminal setting velocity while still travelling in the air as the particle being studied. As shown in the following equation, [109,146]:

\[ daer = dg_{\text{read}} \frac{\rho}{x \cdot \rho_0} \]  

Where daer is the aerodynamic diameter, dg is the geometric diameter, ρ is the particle density, ρ0 is the unit density (usually from water), and x is the shape factor. [147] However, at such a small scale, compared to the extent of the dispersive forces that could be encountered during aerosolization, the particles exhibit strong adhesive and cohesive interparticulate forces, raising the probability of agglomeration. Therefore it is highly vital to fluidize and disperse the micronized particles before they reach the respiratory airways. [148] Based on the dose metering system, DPIs are typically divided into three categories: single-unit dose, multi-unit dose, and multi-dose reservoir.

Dry powder systems use single or blended drugs with an effective carrier, usually lactose, for delivery to the lungs. The three key factors that influence the pulmonary delivery of drugs may be the medication, carrier, and system. In comparison to MDIs, the delivery of drug with a DPI involves minimal patient communication and breathing coordination after the system is actuated. [149] Furthermore, DPIs are lightweight, portable devices which can be held in a purse or pocket easily. Spacers are also not necessary to be used. [123] In fact, as typically needed in MDI formulation, DPIs are absent of environmental-injurious CFC propellants. Since both MDI and DPI were exposed to afford comparative effectiveness in the delivery of the same drug and in view of the United Nations compulsory ban on the use of CFCs in MDIs, it is not surprising that DPIs have become increasingly relevant over the previous decade as a pulmonary drug delivery method. [150] The distribution of aerosol particles drugs is undergoing drastic modifications in all aspects of the inhaler system and formulation. Dry powder inhaler devices are especially attractive. From the other side, because of the possible physical instability of the powder, the formulation and processing of dry powders for inhalation can be difficult and demanding.

Advantages for Dry Powder Inhalers: Relatively large drugs for pay loads per puff are available. Blending is not necessary for this. The particles easily disaggregate despite their small size, enabling the use of small flow-rate independent inhalers. Enables improved pulmonary deposition, reduction of dose uncertainty and compressed dose capacity by improved dispersibility. [151,152]
**Nebulizer**

Today, nebulizers are often used by many doctors to treat acute asthma in an emergency care unit or to treat patients with serious asthma at home. In jet nebulizers, an aerosol is created by directing a high-velocity air stream from a pressurized source against a thin layer of liquid solution. Ultrasonic nebulizers contain the sound of the solvent being aerolized by a piezoelectric crystal. More medications can be delivered to the lungs by the nebulizer than MDI or DPI. The most prevalent disadvantage of the nebulizer is its lack of flexibility, as well as higher drug delivery costs due to the increased demand for healthcare practitioners to assist, and the need for higher drug doses to produce a therapeutic outcome.

**Metered Dose Inhaler (MDI)**

These are the most common instruments for aerosolized drug administration. In this procedure, a drug is combined with a propellant in a canister, and upon actuation of the system, the preformed mixture is expelled in exact calculated quantities. Appropriate use of MDIs allows patients to understand how the system works to organise exhalation and inhalation. Using the spacer system, the issue can be marginally overcome by avoiding the cumbersome size of the device for patients who need to use MDIs outside their houses. Marking the start of 1990, as a result of the mandatory ban on the use of propellant chlorofluorocarbons, which were implicated in the degradation of the Earth’s ozone layer, efforts were deliberately made to reformulate MDIs.

**Particle Engineering Technologies for Pulmonary Delivery**

Researchers have used particle-engineering technologies to change the physicochemical properties of particles, such as decreasing particle size and/or density, changing particle shape and surface characteristics, and changing crystalline morphology, in order to improve particle deaggregation by increasing lung deposition. It is one of the most common and oldest particle processing methods used in the pharmaceutical business. Dry or wet pearl-ball milling, jet milling, or high-pressure homogenization are some of the methods that can be used.

**Spray Drying**

A medicine and excipient solution is sprayed through nozzles into a drying chamber in the spray-drying process, where the solvent is evaporated by a hot-air cyclone. Biodegradable and biocompatible polymers such as poly(lactic-co-glycolic acid) spray-dried polymeric microcarrier systems have allowed the production of several engineered particles loaded with drug micro- and nanoparticles for lung delivery. Spray-drying has been commonly used in dry powder formulations for the pulmonary delivery of proteins such as insulin and viruses.

**Spray-freezing Methods**

An aqueous drug solution is sprayed into a cold vapour over a cryogenic liquid to form droplets in a spray-freeze-drying technology. The droplets are then lyophilized with good flowability, producing micronized and nanosized powders. Using this technology, spherical, highly porous excipient-free particles can be created.

**Thin Film Freezing**

A drug solution containing a bulking cryoprotectant is rapidly frozen dropwise onto a rotating cryogenic substrate in thin film freezing. For solvent removal, the frozen discs are collected and freeze-dried. The engineered particles produced from this process form a brittle powder matrix of low density that is easily dispersed when aerosolized to produce respirable particles from a DPI unit.

**Sono-crystallization**

Ultrasound waves are applied during the drug solution’s antisolvent crystallisation process to monitor the precipitation in this method. Nucleation and crystallisation are caused by ultrasound, increasing reproducibility and uniformity of particle size.

**Fixed-dose Drug Combination**

Fixed-dose drug combination formulations must ensure the homogeneity of the powder and the distribution of a standardised dose to patients, particularly when the ratio of each drug contained in the formulation varies significantly. Particle engineering has been used instead of the simplistic mixing of micronized drugs with coarse carrier particles in an effort to satisfy these criteria and achieve co-deposition of drugs in the lungs. The Global COPD Initiative suggests that bronchodilator combinations with various pathways and length of action therapy should be used to improve the degree of bronchodilation while minimising possible side effects. By avoiding disease progression, exacerbation, and offering symptomatic relief, the main goal is to improve the quality of life. It has been hypothesised that when both drugs co-deposit the target cells, the effectiveness may be due to the synergistic activity of fluticasone and salmeterol. The use of a third part, such as a long-acting muscarinic antagonist, such as tiotropium bromide and ipratropium bromide, is recommended when dual therapy is inadequate to regulate exacerbation and breathlessness.

**Nanoparticles and Biodegradable Polymeric Nanocarriers**

The use of nanoparticles can improve drug lung deposition, increase the velocity of drug dissolution, promote nanoparticles’ epithelial cell internalisation, prolong
drug release in the lung, and allow cell-specific targeted drug delivery or modified drug biological distribution.\[175]\]

In order to minimise the amount of polymer load needed for administration, an effective nanoparticle drug-delivery system should ensure high drug loading capacity. However, lung delivery of discreet nanoparticles is unviable due to low lung deposition, as a consequence of the low inertia of the particles and the tendency for such particles to aggregate irreversibly.\[176]\]

**Conclusions**

On a range of formulations for aerosolized delivery of drugs to the lungs, a substantial body of literature is available; many of these include techniques of particle engineering applied to the powder and a significant number more to the many available applicable delivery devices. A better understanding of the overlap of anatomical, physiological, and pathological factors is required, however, and the way in which they interact with the drug and drug delivery devices' physicochemical factors. Some of these factors have been identified that affect the efficiency of delivery of pulmonary drugs, but the complexities of the overall process have yet to be appreciated. The effect of the variable airway geometry and the airflow speeds through the different regions of the respiratory tract must be addressed to maximise aerosol therapy. It is also important to consider the impact of age disparities and health conditions on airway calibre and airflow patterns. Computational particle deposition simulations can benefit, but they still suffer from restrictions that obstruct their universal use and application.

The lack of an effective animal model that truly mimics drug delivery to humans is perhaps one of the most significant challenges. Animal models are not indicative of the human condition due to changes in the breathing pattern of animals and differences in the branching of airways. Therefore, it is very difficult to extrapolate findings to humans from animal models and much harder to extrapolate these results to young children and elderly adults. Future challenges include the need for more responsive airway flow tests to be established to adapt passive inhalation experiments in animals to those in humans. Lastly, our knowledge of drug absorption processes in the lungs is still relatively poor. Recognizing the expression of different transporters in different regions of the lung can lead to successful delivery of targeted drugs to particular receptors, thereby enhancing therapeutic results. However, there are a few issues remaining for their successful clinical application. Most of all, potential organ toxicity is a major concern in pulmonary medicine.

**Abbreviation**

IPF: Idiopathic Pulmonary Fibrosis
COPD: Chronic Obstructive Pulmonary Fibrosis

**References**

20. Tronde A, Nordén B, Marchner H, Wendel AK, Lennerstås H, Bengtsson UH. Pulmonary absorption rate and bioavailability...
Influence of Particle Size and Particle Deposition of Inhaled Drug in Lung Disease


78. Lipworth BJ. Pharmacokinetics of inhaled drugs. British journal of clinical pharmacology. 1996;42(6):697-705.


90. Wiggins NA. The development of a mathematical approximation technique to determine the mass median aerodynamic diameter (MMAD) and geometric standard deviation (GSD) of drug particles in an inhalation aerosol sprat. Drug development and industrial pharmacy. 1991;17(14):1971-86.


Influence of Particle Size and Particle Deposition of Inhaled Drug in Lung Disease


