IS THERE A FUTURE FOR THE RESPIRATORY DELIVERY OF BIOLOGICS?

In this article, Igor Gonda, PhD, Founder and Chief Executive Officer of Respidex, covers the field of nasal and pulmonary delivery of biologics, considers past failures and successes, and proposes answers to the challenges faced, from biological, biopharmaceutical and commercial standpoints.

The pulmonary and nasal delivery of biologics is intuitively attractive; it is an easy, non-invasive administration route with readily targetable portals – the mouth and the nostrils. On top of that, there is a lot of existing expertise in delivery of drugs both to the lung and to the nose for the treatment of diseases in those parts of the body, such as asthma, COPD, pulmonary arterial hypertension, cystic fibrosis, rhinitis and allergies. These types of therapies continue to attract investment and are growing in use (Figure 1). However, current sales are almost entirely for small molecules. So why is it that there are so few biologics on the market delivered by these routes of administration, even for the treatment of respiratory diseases? How could it change?

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After the heyday of pulmonary and nasal delivery of macromolecules in the 1990s, including proteins, peptides and gene therapies, it is quite remarkable how little progress has been made and how devastating the attrition rate has been for the vast majority of these products before and, in a few cases, after they reached the market.

This article will discuss some of the key factors contributing to the current lack of enthusiasm in this field and provide some thoughts as to what can be done to bring this very promising source of new therapies back into use.

Figure 1: Potential for nasal and pulmonary delivery of biologics.

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therapeutics back on the stage to lead to attractive prophylaxis and treatment for a number of serious diseases.

**BIOLOGICAL BARRIERS TO EFFECTIVE RESPIRATORY DELIVERY**

The respiratory tract is the only part of the body which must be in continuous contact with the environment for us to survive. It is therefore not surprising that, in order for it to continue functioning over the course of many years, it has a multitude of protective mechanisms that act as barriers to entry of foreign materials, as well as other mechanisms that act to remove such materials should they penetrate beyond these barriers. Understanding the nature of these mechanisms and our ability to overcome them, or use them to our advantage, is therefore important.

**Barriers Prior to Deposition**

It goes almost without saying that orally or nasally inhaled substances cannot reach those parts of the respiratory tract that are not ventilated. This is particularly important to consider in the context of obstructions of the respiratory tract, be it "stuffy nose", or the presence of permanent or reversible airway obstructions in asthma and COPD. Combinations of drug, such as decongestants, bronchodilators, mucolytics, and non-drug therapies, such as physiotherapy, prior to biologic delivery, are examples of how some of these reversible obstacles can be overcome.

The human nasopharynx and oropharynx are especially efficient at filtering out large “non-respirable” particles. In the case of the nasopharynx, no matter whether the purpose is the delivery of drugs to the nose for local therapeutic treatment, or via the nose into the systemic circulation or the brain, this highly efficient filter is the desired “barrier” and the target for the therapy.

Delivering large particles, or jets of fluid at high velocities, will maximise nasal deposition – and minimise the possibility of undesirable passage to the airways and beyond. However, the nose is anatomically complex and such a simplistic approach is far from able to achieve optimum targeting of various parts of the nose that may have different functions of therapeutic interest.⁴

A considerable number of studies have been done on preferential targeting to different parts of the nose.⁵,6 For example, delivery of substances via the olfactory nerve into the brain requires delivery to the posterior upper nasal cavity.⁷ Furthermore, it is not always clear what roles different parts of the nasal cavity have in terms of the desirable versus undesirable biological effects. Vaccination via the nose is likely to require targeting the dendritic cells, but their location appears to be mobile and also dependent on the disease and its treatment.⁸

Deposition of orally inhaled therapeutics in the mouth is invariably wasteful, except in relatively rare situations when the oral cavity is the therapeutic target. Because of the complex and dynamic anatomy of oropharynx, deposition there is also the most likely cause of intra- and inter-individual variability in lung delivery. Therefore, minimising deposition in the oropharynx is important for drugs where efficiency and precision of delivery matter. Depending on their aerodynamic properties, particles and droplets that penetrate these initial obstacles will deposit by impaction, interception (in the case of elongated particles), sedimentation and diffusion. If the residence time is insufficient for diffusion and sedimentation of the smallest particles to the walls of the respiratory tract, then they may be exhaled.⁹

There has been extensive research on preferential targeting to different parts of respiratory tract. The key control parameters are the aerodynamic size distribution of the particles carrying the drug, the inspiratory flow rate and inspired volume. For bolus aerosols, placing of the bolus at the beginning of a slow inspiration and then chasing it with clean air provides the maximum probability of deposition in the distal airways and the alveoli. This very large absorptive surface area is ideal for systemic absorption of macromolecules, provided the carrier particles have small enough aerodynamic diameters (<3 μm) to escape inertial deposition in the oropharynx and large airways. Adequate time (approximately 10 s) is required for breath-holding to maximise the deposition of these particles by sedimentation and diffusion. Alternatively, hygroscopic growth of particles was proposed as a mechanism to inhae initially small particles that will then grow by absorbing moisture to maximise their deposition in distal parts of the respiratory tract.⁹

**Post-deposition Mechanisms**

The complexity of delivering therapeutics to the nose is due both to its anatomy and multiple mechanisms affecting the transport of drugs through it. Even the same anatomical region may handle particles deposited in adjacent places differently. For large molecules, absorption from the nose is relatively slow while the aforementioned mechanisms are quite fast. Therefore, to maximise the therapeutic activity in the nose, or to achieve efficient absorption into the systemic circulation, the means to increase the residence time in the nose and to prevent unproductive elimination needs to be proactively sought. Most efforts in this area have involved the use of bio-adhesive drug carriers, and enzyme inhibitors.

The nasal cavity has extensive intra- and extra-cellular enzymatic activity, containing both peptidases and proteases. Studies with peptides showed that, by using peptide inhibitors, it is possible to enhance nasal absorption. Indeed, to improve the bioavailability of nasally delivered therapeutics of this nature, enzyme inhibitors paired together with formulations that increase the residence time in the nose may be necessary. Otherwise, even for relatively small peptides, the systemic bioavailability is of the order of approximately 1%. Biologics that are resistant to enzymatic degradation in the nose are an alternative to the use of enzyme inhibitors.

**Deposition Of Biologics in the Airways and Alveoli**

Once particles deposit on conducting airways, mucociliary clearance will start translocating them upwards and they may ultimately get swallowed. Macrophages, and possibly other phagocytic cells, can
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also ingest such particles which can be used to our advantage if they are the therapeutic target.

The lung has relatively low metabolic activity compared with the gastro-intestinal tract. The presence of protease activity is generally associated with lung diseases, such as alpha-1 antitrypsin (A1AT) deficiency with absence or abnormally low levels of the natural protease neutraliser A1AT, or in cystic fibrosis, where high levels of proteases are the result of chronic inflammation and infection.

There is evidence from animal studies that peptidase inhibitors increase the bioavailability of inhaled peptides such as insulin and calcitonin delivered to the lung.15,24 Mucociliary clearance is absent from alveoli but alveolar macrophages can phagocytose particles there.

These removal mechanisms compete against the biologics’ intended therapeutic activities, be it binding to receptors within the respiratory tract, or absorption. Protection against metabolic activity can be achieved by encapsulation in carriers. Liposomes have the advantage of resembling the natural components of the lung and have been extensively tested in large late-stage clinical trials with small molecules.15,16

MARKET BARRIERS TO DELIVERY OF BIOLOGICS

Cost
Most biologics are much more expensive to manufacture than small molecules. The low systemic bioavailability of the nasal and pulmonary routes requires much higher doses than injections, which for costly biologics may render these administration modes less attractive.

In contrast, delivering biologics directly to the sites of therapeutic action in the respiratory tract makes these routes very attractive not just from the cost-of-goods perspective but also to reduce the potential for systemic side effects. An example of such a product is recombinant human DNase for the treatment of cystic fibrosis – one of the few approved inhaled biologics to have achieved sustained success.

Safety
There have been relatively few studies published on the results of long-term animal and human safety of biologics delivered by inhalation. The concern about carcinogenicity of inhaled insulin expressed by Pfizer with respect to their inhaled product Exubera®, which they dropped and returned to Nektar Therapeutics (San Francisco, CA, US),17 undoubtedly adversely affected not only other inhaled insulin products, but also the whole field of delivery of biologics by inhalation. Although the number of lung cancer cases associated with the use of Exubera (all in smokers) and MannKind’s Afrezza® (smokers and non-smokers) has been low, it remains a concern.18,19 To the best of my knowledge, no other biologics have reported carcinogenicity findings during their development or post-approval. The most extensive published animal and human safety studies appear to be for recombinant human DNase.20 Studies with inhaled proteins suggest generally good safety.21

Lackluster Performance of Marketed Inhaled Insulin Products
Unfortunately, the widespread exuberance about the market potential for Exubera, the first approved inhaled insulin product, followed by failure to achieve even a small fraction of the sales forecasts had a massive negative impact on the whole field of inhaled delivery of biologics. And, in more recent times Afrezza, the second approved inhaled product, which overcame several of the weaknesses of Exubera (e.g. large inhaler, complicated instructions for use and maintenance) also fell far short of expectations.

There is no single explanation for the commercial disappointments with these products.18 The common factors for both Afrezza and Exubera were:

• Fast-acting insulin only. Since many diabetics need to use both short- and long-acting insulins, neither of these products provided a full solution for patients. Future products of this kind need to look more holistically to solve the patients’ needs.

• The lack of experience. Patients and their caregivers, as well as the sales and marketing forces behind the products, lacked experience with inhaled products, including the requirements for regular respiratory safety testing and future efforts in this area will require better education of all key stakeholders. First and foremost, the development scientists have to design the new product such that it meets the patients’ needs and is more attractive compared with existing therapies, reducing rather than increasing the burden on the patient and the provider.

• Significant improvement of injectors. During the development of Exubera and Afrezza insulin, injection device technology saw notable advancement. This is a tough area to deal with, but anticipation of evolutionary improvements in existing products, such as smaller gauge needles, that could then become highly competitive against a revolutionary product, in this case inhaled insulin, before and during development and marketing the product could save a lot of later disappointment.

• Both products were dry powders, with evidence for respiratory side-effects. Patients with asthma and COPD were excluded from the labels of both Exubera and Afrezza. Yet, apart from hypoglycaemia, the most common side-effects were cough and throat pain or irritation. Using formulations and devices that minimise upper and central airway deposition, with compositions that have a low probability of airway irritation, may overcome these issues and possibly improve the uptake of such products. An example of such development was AerX iDMS that used an aqueous formulation of insulin delivered with a highly efficient inhaler (Aradigm Corporation, Hayward, CA, US) targeting alveoli with small, nearly monodisperse droplets delivered as a bolus at the beginning of a slow inhalation. These formulations were well tolerated even in patients with asthma and COPD. Whether such attributes will significantly improve market penetration through better tolerability and ability to include patients with such common co-morbidities remains to be seen.
Slow Entry of Biosimilars

One of the commercial drivers for research into the delivery of biologics via pulmonary and nasal delivery was the anticipation of rapid introduction of biosimilar copies of approved biologics. It was expected that, as with small molecules, changes in the route of administration from injections to more convenient, non-invasive methods would provide patient-attractive competitive advantages to the company introducing such a product, as well as the possibility of significant extension of exclusivity through delivery-related patents.

However, the path to biosimilar approvals turned out to be much more complex than for small molecules. In the US, identical copies of existing drug products, i.e. generics, can be approved on the basis of a very much abbreviated approval process, the US FDA’s ANDA. Innovative improvements of already approved small-molecule drugs, such as changing the route of administration from injections to oral or nasal inhalation, can be pursued via the 505(b2) regulatory path that can utilise the existing public information about the drug’s safety and efficacy. Such products are often patent-protected and may also gain regulatory exclusivity via other means, such as via an orphan drug designation.

Because of the much increased complexity of biologics, the regulatory rules for biosimilars have taken a long time to evolve and implement. These factors have contributed to a relatively slow entry of such products, especially into the US market. Consequently, neither the originators of the products nor the new entrants have embraced alternative means of delivery, including nasal and pulmonary, to make more competitive “bio-betters”. Considering the risk and cost involved with the development of new therapies based on new molecules, and especially on new mechanisms of action, it is possible that bio-betters based on more attractive delivery methods will be pursued now.

There is already much concern about the relative cost/benefit of the recent market entries of biologics for the treatment of asthma. This is an obvious area where inhalation products using the same, or similar molecules, delivered directly to the site of action in the respiratory tract, are likely to improve the therapeutic ratio (efficacy/safety) and potentially reduce the cost of raw materials.

Fear of the Unknown

Development of new products and technologies using pulmonary and nasal routes for biologics is not mainstream pharmaceutical and biotech activity, as the default option remains injectables. Only when it is recognised that locally administered treatment is practically inevitable, as in the case of recombinant human DNase and gene therapies for cystic fibrosis, will these routes be taken to be the first option.

I have a very vivid memory from sitting on a panel of experts on delivery of biologics in the early 2000s at an investors’ conference; a senior executive from a large biotech company seriously claimed that pulmonary delivery of biologics was “impossible”. Yet, by that time recombinant human DNase was already approved and successful results with inhaled biologics in humans were published in reputable journals. I would not be surprised if that is still a widespread perception outside the close-knit community of respiratory delivery aficionados.

This is not helped by the fact that the causes for some of the very disappointing failures in late stage clinical trials with inhaled biologics were never explained, such as the inhaled gene therapies and Alpha-1 Antitrypsin, or lung surfactant therapies. On the other hand, the low efficacy of recombinant human DNase outside cystic fibrosis is quite well understood and using improved molecules and better prospectively identified patients may make modified recombinant human DNase viable in indications such as COPD.

CONCLUSION

Delivery of several types of biologics via the pulmonary and nasal routes is clearly feasible and may lead to very valuable products for patients whose needs are not met today, or only currently met with a high treatment burden. Maximising efficiency through the improved delivery of biologic agents to the desired sites and their protection against unproductive elimination is desirable, and there is both extensive expertise and availability of much improved devices and formulations now to help achieve these goals.

Previous approvals of injected biologics, taken together with experience with approval processes of current pulmonary and nasal products, will certainly assist in shaping the regulatory paths for these new products. Since it is likely that the control groups in clinical trials will be patients using the existing approved route of administration for the same biologic, enrolment for clinical trials as well as choice of endpoints should be much easier and the risk much lower than for placebo-controlled trials of new drugs or biologics.

But technical optimisation and abbreviated regulatory paths alone will not be a guarantee for success. The development teams working on the next generation of inhaled biologics, especially for those to be used outside respiratory medicine, should learn from...
history; inhaled products should be designed, developed and marketed through close collaboration with the patients and their healthcare providers to be attractive for all key stakeholders. Collecting information during development to support the socio-economic benefits of these new products is important, beyond the need to prove efficacy and safety.

ABOUT THE COMPANY

Respidex is a consultancy helping pharmaceutical companies in their R&D activities, regulatory strategy, financing and commercialisation of products for prevention, diagnosis and treatment of diseases. Respidex clients include small early preclinical companies as well as enterprises in late stage clinical development.

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ABOUT THE AUTHOR

Igor Gonda, PhD, is the Founder of Respidex, a San Francisco-based consulting company. Dr Gonda has held a variety of executive positions in US and Australian private and public pharmaceutical and biotech companies. His research focus was primarily in the area of drugs and biologics administered to the lung and nose conducted in industry and, prior to that, in universities in England, Australia and the US. Dr Gonda’s current interests additionally include the involvement of patient advocacy groups in pharmaceutical R&D, venture philanthropy and the regulatory process to accelerate availability of cures for their diseases, and in opportunities for making individualised precision medicines faster and cheaper.