Chapter 17

Bacteriophage Encapsulation Using Spray Drying for Phage Therapy

Danish J. Malik

Chemical Engineering Department, Loughborough University, Loughborough LE11 3TU, United Kingdom

Email: d.j.malik@lboro.ac.uk

DOI: https://doi.org/10.21775/9781913652517.17

Abstract
Exploiting the potential of bacteriophages for phage therapy is an exciting future prospect. However, in order to be successful, there is a pressing need for the manufacture of safe and efficacious phage drug products to treat patients. Scalable manufacture of phage biologics as a stable solid dry powder form is highly desirable and achievable using the process of spray drying. Spray drying of purified phage suspensions formulated with suitable excipients can be carried out in a single step with high process throughput and at relatively low cost. The resulting phage-containing powders can possess good storage shelf-life. The process allows control over the final phage dose in the powder and production of microparticles suitable for a variety of therapeutic uses. Spray dried powders may include different polymer formulations employing a multitude of different triggers for phage release at the target site including pH, enzymes, virulence factors etc. The activity of the phages in spray dried powders is adversely affected during spray drying due to desiccation and thermal stresses which need to be controlled. The choice of polymers, excipients and moisture content of the dry powders affects the material glass transition temperature and the stability of the phages during storage. The storage temperature and storage humidity are
important factors affecting the stability of the phages in the dry powders. A quality by design (QbD) approach for phage drug product development needs to identify drug product characteristics that are critical to quality from the patient’s perspective and translates them into the critical quality attributes (CQA) of the drug product. The relationship between the phage drug product CQAs and formulation development and spray drying process conditions are discussed in this chapter.

**Introduction**

Phage drug products need careful formulation development and an appreciation of the chemical and physical stresses that bacteriophages (phages) may encounter both during processing as well as during storage once formulated. Phage inactivation and long-term reduction in phage titres upon storage are highly undesirable. Delivery of high titres of phages and their controlled release at the site of treatment affects phage pharmacokinetics (Malik et al., 2017). The physical and chemical properties of the materials used in the formulation need careful consideration when selecting a technique for encapsulation. Many techniques and processes may be used for stabilising, immobilising and encapsulating phages including spray-drying, spray freeze drying, freeze drying, extrusion dripping methods, emulsion and polymerisation techniques. (Malik et al., 2017). Phages are protein structures, and they are therefore susceptible to factors known to denature proteins; and these include exposure to organic solvents (Lee and Belcher, 2004; Puapermpoonsiri et al., 2009), high temperatures (Briers et al., 2008), pH (Briers et al., 2008; Knezevic et al., 2011), ionic strength (Knezevic et al., 2011), and, interfacial effects. Additionally, mechanical stresses during formulation or encapsulation including shear stresses during mixing and agitation, atomisation during spraying (Leung et al., 2016) and, desiccation stresses during drying (Dini and de Urraza, 2013), need careful consideration. Scalable, low cost GMP compliant processes for phage encapsulation are needed.

**Spray drying of phages**

Spray drying processes atomise a liquid containing dissolved solids, converting it into a fine mist or aerosol which is contacted with a hot dry gas (typically hot
Dry air) inside a drying chamber (Figure 1). Due to the high surface area to volume ratio of the very small atomised droplets (droplet size typically 10 – 100 µm), the solvent rapidly evaporates with each droplet forming a particle comprising the non-volatile components (solids fraction in feed containing phage and excipients) along with a small amount of residual moisture. The particles are separated from the airstream exiting the dryer, and cyclones or bag filters are typically used for this. Spray drying is a scalable industrial process technology and is widely used to produce fine powders containing biologicals for pharmaceutical applications including pulmonary delivery via dry powder inhalers (DPI) (Hoe et al., 2014) (Figure 2). Dry powders are increasingly being considered for phage drug products because they show relatively long storage stability without requiring a cold supply chain and refrigeration (Carrigy et al., 2019; Chew and Chan, 2002; Klingler et al., 2009). Spray drying phages using formulations containing synthetic polymers may allow encapsulation, targeted delivery and controlled release aspects such as triggered release in response to pH changes, e.g. for gastrointestinal delivery (Vinner et al., 2019). The powders may also be filled into capsules or compressed into tablets for oral solid dosage (Vinner et al., 2019).

![Figure 1. Schematic diagram illustrating a typical spray drying set-up to produce phage powders.](image-url)
Spray drying of phage biologicals using a Quality-by-Design approach

Quality by design is a concept pioneered by Joseph Juran and is based around the concept that quality should be designed into a product rather than relying on end-of-pipe testing (Juran, 2008). A high-quality phage drug product needs to be safe and efficacious, delivering the therapeutic benefit promised to the consumer on the product label. Regulators such as the US Food and Drug Administration (US FDA) encourage manufacturers to adopt a risk-based
approach using QbD principles for drug product development, manufacturing and regulation (Yu et al., 2014). A QbD-led approach emphasises a deep understanding of the manufacturing process and linking product quality to the desired clinical performance of the drug product. Designing a robust formulation and manufacturing process for phage drug products that consistently deliver the desired phage drug product quality is essential for the future of phage therapy. Performance-based quality specifications are critical in ensuring this is achieved. Improved product and process understanding helps in designing control strategies to ensure the drug product quality remains within acceptable limits without compromising safety and efficacy. Spray drying of phage biologicals requires identification of process and product characteristics that are critical to product quality from the patient’s perspective. These then need to be translated into critical quality attributes (CQAs) of the phage drug product. Linking formulation and manufacturing aspects to CQAs is vital in ensuring the phage drug product meets performance quality criteria. This requires defining what the phage biological’s quality target product profile (QTPP) looks like and linking it to CQAs. Identification of critical material attributes (CMAs) (explained below) that are relevant to spray drying of phage biologicals needs to be ascertained as well as critical process parameters (CPPs) specific for the spray drying process. Implementation of process control strategies and accounting for variability in process inputs as well as putting specifications on CMAs and the drug substance, excipients and drug products are needed (Yu et al., 2014).

The QTPP of the phage drug product is the basis for the phage drug product development and is a summary of all the quality attributes that the drug product needs to meet to ensure it is safe and efficacious (Yu et al., 2014). Identification of the phage drug product’s CQAs is the next step and includes many attributes including phage titre in the powder, the powder moisture content, the material’s glass transition temperature etc (see summary Table 1). The QTPP of new phage drug products needs to be adequately defined before drug product development work commences. A focus on the phage product design is just as important as the consideration of the process design here. The stability of the phage drug product throughout its shelf life needs to be confirmed through stability studies. The key objective of product design is to ensure that the phage
drug product retains the desired QTPP over the product shelf life (Yu et al., 2014). Failure of a recent phage therapy trial due to a complex liquid phage drug product having stability issues during the trial is a valuable lesson for the future (Jault et al., 2018). Key aspects related to CMA and CQA for dry powder phage biologicals relevant to spray drying of phage formulations are discussed below.

Table 1. Typical input material attributes, process parameters and quality attributes of spray dried phage powders.

<table>
<thead>
<tr>
<th>Input material attributes</th>
<th>Process parameters</th>
<th>Quality attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Phage type, e.g. <em>Myoviridae</em></td>
<td>• Drying temperature</td>
<td>• Phage titre in spray dried powders (potency)</td>
</tr>
<tr>
<td>• Intrinsic susceptibility of a phage drug candidate to spray drying process stresses including atomisation, drying, desiccation, osmotic and re-hydration stresses</td>
<td>• Drying air gas flow rate</td>
<td>• Powder glass transition temperature (T_g)</td>
</tr>
<tr>
<td>• Excipient type, e.g. sugars, polymers, proteins</td>
<td>• Liquid feed flowrate</td>
<td>• Powder moisture content</td>
</tr>
<tr>
<td>• Solution viscosity</td>
<td>• Type of atomiser, e.g. two fluid nozzle (and nozzle size), rotary wheel (and rotation speed).</td>
<td>• Phage shelf life (Months stored under storage specific conditions at a given temperature and relative humidity)</td>
</tr>
<tr>
<td>• Solution solids content w/v%</td>
<td>• Dryer residence time</td>
<td>• Powder re-suspension characteristics</td>
</tr>
<tr>
<td>• Phage titre</td>
<td>• Cyclone collection efficiency (affects the yield of material)</td>
<td>• Powder particle size and size distribution</td>
</tr>
<tr>
<td>• Lysate purity</td>
<td></td>
<td>• Cohesive/adhesive properties</td>
</tr>
</tbody>
</table>

Malik  Bacteriophage Encapsulation
The role of excipients in stabilising spray dried phage drug products

Spray drying of phage suspensions typically includes excipients in the formulation to protect the phages from thermal and desiccation stresses. The critical material attributes (CMAs) of excipients used in spray drying needs careful consideration. Trehalose, sucrose and lactose have emerged as the most promising excipients for spray drying phage suspensions with trehalose being the front runner due to its high glass transition temperature and water replacement properties (Grasmeijer et al., 2013). Trehalose is the most frequently reported excipient used in the spray drying of phages and it has been shown to result in spray dried powders with high phage titre and good storage stability (Carrigy et al., 2019; Leung et al., 2016, 2017; Matinkhoo et al., 2011; Vandenheuvel et al., 2014). Upon spray drying, sugar excipients form amorphous structures with high glass transition temperatures (Tg), e.g. in their anhydrous amorphous form Tg of trehalose is 115 °C and Tg for lactose is 108 °C (Chang et al., 2019; Chen et al., 2000; Roe and Labuza, 2005). Below the glass transition temperature, vitrification due to the excipient stabilises the encapsulated bioactive agent and permits storage over long periods at ambient or lower temperatures (Carpenter and Crowe, 1988; Leung et al., 2017). Trehalose has low toxicity and has been shown to protect biological materials including proteins, probiotics and vaccines, against desiccation and thermal stress. Crowe et al. (1996) suggested that the efficacy of trehalose as an excipient in drying is partly due to its high glass transition temperature at all water contents but also because trehalose binds residual water left over from the drying process to form a dihydrate which might otherwise participate in lowering the glass transition temperature to below ambient (Vandenheuvel et al., 2014). In the amorphous state, trehalose may stabilise protein conformation through hydrogen bonding (Allison et al., 1999). Other excipients that have been used to improve the dispersibility of spray dried phage containing powders include dextran, lactose (common excipient used for DPI powders), glucose, sucrose, mannitol and leucine (Leung et al., 2016; Matinkhoo et al., 2011; Telko and Hickey, 2005; Vandenheuvel et al., 2013b). Reducing sugars such as lactose may be unsuitable excipients for phage (Vandenheuvel et al., 2013b). Alternatives such as mannitol have been suggested in the literature; however,
mannitol has a tendency to recrystallize rather than remaining amorphous (Steckel and Bolzen, 2004). Phage spray drying studies using mannitol have reported improved particle dispersion characteristics. However, phage viability results have been mixed (Leung et al., 2016, 2017). Addition of proteins, e.g. casein in combination with trehalose has shown good results (Matinkhoo et al., 2011).

**Effect of spray drying conditions on phage viability and shelf life**

The spray drying conditions need to be carefully optimised to ensure viability of spray dried phages in the final drug product. Two-fluid nozzles typically used for atomisation of the phage suspensions for spray drying have been shown to generate shear induced damage and loss of phage titre (Leung et al., 2016; Vandenheuvel et al., 2013b). Phages are highly sensitive to thermal stresses resulting in loss of phage activity at high spray drying temperatures (for co-current dryers, outlet temperatures exceeding 60°C) (Davies and Kelly, 1969; Jończyk et al., 2011). Typically, spray drying studies with phages have employed co-current dryers operating at low outlet spray drying temperatures (~40-50 °C outlet air temperatures) resulting in higher phage titres in the powders and reduced overall phage losses immediately post spray drying (Leung et al., 2017). High loss of phage titres have been observed at higher spray drying temperatures (Leung et al., 2016; Matinkhoo et al., 2011; Vandenheuvel et al., 2013b; Walbeck, 2013). However, the residual moisture content in the spray dried powders (typically ~5% w/w or more, Figure 3) due to low drying temperatures (40-50 °C dryer outlet temperatures) may result in lowering of the glass transition temperature (Tg) (Roe and Labuza, 2005). The glass transition temperature of an amorphous pharmaceutical solid is a critical physical property (CQA) which can dramatically influence its chemical and physical stability as well as its viscoelastic properties (Figure 4). Water acts as a plasticiser in spray dried powders which may also adsorb water upon storage due to exposure to a humid atmosphere (Hancock and Zografi, 1994). The Tg of amorphous powders such as trehalose, lactose and sucrose fall dramatically when the powders contain low amounts of moisture (Chen et al., 2000; Hancock and Zografi, 1994; Roe and Labuza, 2005). Spray dried phage containing powders tend to be amorphous upon production; however, storage conditions may subject the
material to temperatures and humidity conditions lying above the glass transition line. Crystallization rates of glasses depend on the storage temperature in relation to Tg. The recrystallisation of amorphous sugar in spray dried powders negatively affect phage viability, perhaps due to denaturing of phage receptor proteins as sugar molecules can no longer interact through hydrogen bonding with proteins to retain protein conformation upon crystallisation (Carpenter and Crowe, 1988; Roe and Labuza, 2005) (Figures 5 and 6). The use of excipients that are hygroscopic, e.g. lactose and trehalose also require careful handling under controlled atmospheric conditions and the use of a dry atmosphere controlled glove box is advisable during powder handling to prevent exposure of the dry powder to moisture in the atmosphere (Carrigy et al., 2019; Leung et al., 2017; Matinkhoo et al., 2011).

Figure 3. Measurement of residual moisture in spray dried powders using thermal gravimetric analysis (TGA). Spray dried powder produced at an outlet temperature of 65 °C.
**Figure 4.** Measurement of glass transition temperature using differential scanning calorimetry (DSC) of a spray dried powder (using trehalose as excipient). The powder was produced at an outlet temperature of 65 °C. Onset temperature shown is 50 °C.

**Figure 5.** Measurement of glass transition temperature using differential scanning calorimetry (DSC) of a spray dried powder (using trehalose as excipient). The powder was produced at an outlet temperature of 65 °C and exposed to storage conditions of 30 °C and 65% relative humidity for 24 hours. Peaks indicate re-crystallisation of the trehalose with endothermic melting peaks for different trehalose polymorphs shown here.
Figure 6. Top Panel: Powder X-ray diffraction results showing amorphous trehalose powders immediately, six months and one year following spray drying at 65°C outlet drying temperature and stored at 4°C (blue curve (immediately after spray drying), green curve (six months after spray drying) and red curve (one year after spray drying)). Bottom panel: Same sample exposed to storage conditions of 30°C and 65 % relative humidity for 24 hours resulting in re-crystallisation of the trehalose.

Effect of storage conditions on phage stability in spray dried powders
Storage temperature and relative humidity are the two main factors shown to affect phage stability in spray dried powders. Amorphous forms of dry powder pharmaceutical biologics are most stable when molecular motion in the disordered state is retarded over a meaningful pharmaceutical timeframe (Hancock and Zografi, 1994). The storage temperature needs to be
considerably below the amorphous powder material’s Tg to slow down molecular motion such that chemical and physical instability of the drug product does not occur over the lifetime of the pharmaceutical product, which is typically in the order of a few years (Hancock and Zografi, 1994). Storage temperatures 50°C below the glass transition temperature are thought to significantly reduce molecular movement and help in lengthening phage powder storage shelf life (Chang et al., 2020; Hancock et al., 1995). Refrigerated storage of spray dried powders at relative humidity exceeding 20% has been shown to result in loss of phage titre whereas samples stored at low humidity were stable (Leung et al., 2017; Vandenheuvel et al., 2014). Disaccharides such as trehalose, sucrose and lactose preserve the bioactivity of phages in the dry state. Refrigerated storage of spray dried powders was reported to yield higher titres of phage during long term storage compared with storage under ambient conditions (Leung et al., 2017). Different phage strains formulated and spray dried under identical conditions showed significant differences in the resulting phage titre suggesting the need for individually formulating each phage to be used in a phage cocktail (Vandenheuvel et al., 2013b, 2014). The selection of phages as suitable candidates for phage therapy should include consideration of whether they lend themselves to spray drying i.e. a CMA as part of QbD approach to linking drug product design to the QTPP if the final drug product is to be delivered to patients in a stable dry powder form. A number of studies have reported spray dried phage powders to have a suitable mass median aerosolized diameter for pulmonary delivery of phages to treat respiratory infections (Leung et al., 2016; Matinkhoo et al., 2011; Vandenheuvel et al., 2013a, 2014). Here the powder particle size is a CQA linking powder aerosolization to QTPP. Such critical issues need greater consideration early in the product design cycle.

Conclusion
Formulation and stability of phage drug products need greater consideration during phage drug product design. Spray drying is a scalable, high throughput process that lends itself to the production of stable dry powder phage biologicals. A QbD approach to phage drug product design is advocated in this chapter. CMAs includes the selection of suitable phage candidates at an early
stage in drug product design that lend themselves to spray drying and the use of excipients that result in dry powders suitable for phage biologicals. A better understanding of the critical process parameters related to spray drying and consideration of powder handling, packaging and storage conditions is needed. Spray drying of phage biologicals can increase the stability and shelf life of the drug product as well as incorporating targeted delivery and controlled release elements. Spray dried powders can be loaded into capsules or made into tablets using direct compression processes giving added flexibility for ease of use.

References


