3D printing of medicinal products and the challenge of personalized therapy

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Abstract

By 3D printing, solid objects of any shape are fabricated through layer-by-layer addition of materials based on a digital model. At present, such a technique is broadly exploited in many industrial fields because of major advantages in terms of reduced times and costs of development and production. In the biomedical and pharmaceutical domains, the interest in 3D printing is growing in step with the needs of personalized medicine. Printed scaffolds and prostheses have partly replaced medical devices produced by more established techniques and, more recently, 3D printing has been proposed for the manufacturing of drug products. Notably, the availability of patient-tailored pharmaceuticals would be of utmost importance for children, elderly subjects, poor and high metabolizers and individuals undergoing multiple drug treatments. 3D printing encompasses a range of differing techniques, each involving advantages and open issues. Particularly, solidification of powder, extrusion and stereolithography have been applied to the manufacturing of drug products. The main challenge to their exploitation for personalized pharmacological therapy is likely to be related to the regulatory issues involved and to implementation of production models that may allow to efficiently turn the therapeutic needs of individual patients into small batches of appropriate drug products meeting preset quality requirements.

Keywords: drug delivery systems, solid dosage form, extrusion, oral drug delivery, controlled release, 3D printing, personalized medicine, solidification of powder, fused deposition modeling, stereolithography
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Abbreviations:

2D, two dimensional

3D, three dimensional

ABS, acrylonitrile butadiene styrene

API, active pharmaceutical ingredient

ASTM, American Society for Testing Materials

CAD, computer-aided design

DDS, drug delivery system

DPPO, diphenyl (2,4,6-trimethylbenzoyl) phosphine oxide

EC, ethyl cellulose

EDTA, ethylenediaminetetraacetic acid

EVA, ethylene vinyl acetate

FDA, Food and Drug Administration

FDM, fused deposition modeling
GRAS, generally recognized as safe
HIPS, high impact polystyrene
HME, hot melt extrusion
HPC, hydroxypropyl cellulose
HPMC, hydroxypropyl methyl cellulose
IM, injection molding
MCC, microcrystalline cellulose
PCL, poly-ε-caprolactone
PEG, polyethylene glycol
PEGDA, poly (ethylene glycol) diacrylate
PLA, poly (lactic acid)
PLGA, poly(lactide-co-glycolide)
PVA, polyvinyl alcohol
PVP, polyvinyl pyrrolidone
SLA, stereolithography
SLS, sodium lauryl sulfate
TEC, triethyl citrate
TPU, thermoplastic polyurethane
UV, ultraviolet
3D printing glossary:

- **ADDITIVE MANUFACTURING**: the process of joining materials to fabricate objects from 3D model data, usually layer by layer

- **COMPUTER-AIDED DESIGN**: refers to the use of computer systems to assist in the creation, modification or optimization of 3D drawings

- **EXTRUSION**: 3D printing process that involves layer by layer deposition of molten/softened or liquid/semisolid materials through a syringe or a nozzle

- **FUSED DEPOSITION MODELING**: 3D printing technique that involves the use of one or more filaments, generally based on a thermoplastic material, which are extruded and deposited in a molten/softened state layer by layer

- **RAPID PROTOTYPING**: production of a prototype/scale model of a physical object/assembly using 3D computer-aided design data

- **RESOLUTION**: degree of conformity of a 3D printed object to the electronic model by which it was generated

- **SOLIDIFICATION OF POWDER**: 3D printing technique that involves the distribution of thin layers of powder selectively joined by using drops of a liquid binder

- **STEREOLITOGRAPHY**: 3D printing technique that involves the use of a UV laser to cure a photo-reactive resin, contained in a vat, layer by layer into a solid object

- **.stl FILE**: file generated by a computer-aided design program that contains all the data describing the layout of a 3D object and it is the most commonly used file format for 3D printing

- **SUBTRACTIONAL MANUFACTURING**: process in which 3D objects are fabricated by successively cutting material away from a solid block of the latter
1. Introduction

3D printing enables the construction of solid objects of any shape through layer-by-layer addition of materials (additive manufacturing) based on a digital model\textsuperscript{1,2}. In contrast to traditional subtractive manufacturing, which consists in successively cutting away material from a starting block in order to build up the final object, 3D printing allows production to be carried out with no waste of materials and no need for their disposal. Because objects are constructed automatically, according to preset digital models, relevant preliminary geometry study, production planning and manual manufacturing steps can be limited or avoided. This results in major advantages in terms of reduced times and costs, irrespective of the extent of complexity of the item shape. The degree of conformity of an object to the electronic model by which it was generated (\textit{i.e.} resolution) basically depends on the thickness of the single layers deposited, and can thus be enhanced through proper modulation of this parameter.

3D printing encompasses a range of techniques differing from each other in the nature of the substrate (\textit{e.g.} ceramics, metals, polymers, composites), deposition mode, mechanism of layer formation, printer employed and characteristics of the final product (\textit{e.g.} morphology, texture, surface, thermal/mechanical/conductivity properties). According to the ASTM, these techniques are classified based on the additive process involved (Table 1)\textsuperscript{3}.

Independent of the specific technique applied, 3D printing processes generally entail common steps: the creation of a CAD file, its conversion into a .\textit{stl} file, which will be transferred to the equipment, the printer set-up and proper fabrication of the object, its removal from the build plate, post-processes and final cleaning step if needed (Figure 1)\textsuperscript{1,2}. The entire manufacturing process occurs as a single step under computer control, thus avoiding intermediate stages of production and any manual task. Any change in the final object design can be achieved by modifying the relevant CAD file.
2. Personalization

The current popularity of 3D printing in various fields is often described as a new industrial revolution\textsuperscript{4,5}. While in the last centuries industries were focused on mass-manufacturing, automation and standardization in order to reduce costs and increase profit, with the advent of 3D printing they are shifting to on-demand production of either a small number of objects or even one object at a time (small batches), possibly customized, at affordable prices. Products characterized by complex geometries can also be fabricated and real-time modified to meet individual needs at little or no extra costs. 3D printing is spreading throughout all manufacturing stages of production, from the prototyping step to the fabrication of consumer products. By shortening the design, manufacturing and production cycle, thus simplifying the manufacturing chain, this new technology has brought the site of object fabrication closer to that of demand. Moreover, it has proved effective as a rapid prototyping tool useful to evaluate the form, fit and function of many objects before their production on a large scale. Finally, because items are fabricated upon request, waste and inventory can be reduced, and issues related to overproduction are circumvented.

2.1 Personalized medicine

Within the healthcare field, 3D printing has grown in step with the concept of personalized medicine, which has been attracting more and more attention in the last\textsuperscript{6-10}. Personalized medicine generally consists in tailoring medical treatments to the characteristics, needs and preferences of each single patient, and it involves purposely run diagnosis, therapy and follow-up. The concept could be extended to include pre-emptive medicine aimed at reducing the risk of diseases a subject has shown susceptible to, by changing his lifestyle, diet and habits and by advising him on the use of peculiar supplements or drugs. Personalized medicine is not actually a new idea: physicians have noticed over time that patients with similar symptoms may have different illnesses, and that medical treatments may work in some subjects while not being as much effective in others apparently suffering from the same disease. Recent advances in various medical and biomedical fields, from
genomics to imaging, have already allowed patients to be treated and monitored more properly, in closer and closer agreement with their individual needs. By way of example, genetic tests are employed prior to diagnosis, thereby enabling earlier intervention, more efficient drug selection and increased safety. Indeed, the goal of personalized medicine is to drive clinical decision-making by distinguishing in advance those patients who are most likely to benefit from a given treatment from those who would incur costs and side effects without gaining equivalent benefits. In this respect, 3D printing could provide the answer and instrument for moving from mass-production based on the one-size-fits-all approach to manufacturing of small batches of individually developed products that can be modified in real time and possibly fabricated at the point of care\textsuperscript{11-14}.

2.2 Personalized therapy

In the medical field, 3D printing was initially employed by surgeons as an aid in creating 3D models of patients to better visualize their anatomy, particularly in the case of individuals with unique structures or anomalies, which would require complex surgeries\textsuperscript{15-16}. With respect to tissue engineering, medical devices, especially scaffolds and prostheses, currently represent one of the most interesting manufacturing applications of 3D printing\textsuperscript{17}. This technique has partially replaced more established ones (\textit{e.g.} solvent casting and particulate leaching, membrane lamination, molding) since it enables fabrication, starting from biocompatible materials, of items perfectly fitting the anatomical characteristics of the patient as highlighted by diagnostic imaging tools (\textit{e.g.} X-ray, computed tomography, nuclear magnetic resonance). Scaffolds are intended for different functions, such as space filling, 3D structures that organize proliferating cells into a desired tissue, \textit{in situ} vehicles for the delivery of active molecules (\textit{e.g.} antibiotic or anti-inflammatory drugs, growth factors). The ability to fabricate a 3D structure characterized by pre-determined pore dimensions, distribution and interconnectivity was proved to be critical for cell adhesion and growth. Because 3D printing yields finished items with a high-resolution microstructure, it turns out
especially promising in this area. Only in the new millennium, 3D printing was proposed and exploited for the manufacturing of drug products. The availability of custom-made pharmaceuticals, conceived to meet the individual needs of a single patient, would be of special interest for instance to personalize the dose of the conveyed drug, enable co-administration of drugs in multi-therapies, avoid the use of specific excipients involved in intolerances, such as lactose or sucrose, change the flavor or ease. Particularly, patient-tailored products could be of utmost importance for children, elderly subjects, poor and high metabolizers and individuals who are taking multiple medicines that may interact with each other. In the field of personalized medication, sustainability, patentability and ability to produce at lower costs are only some of the several advantages that could justify the interest in 3D printing. Therapy personalization itself was identified as a tool for reducing the healthcare system expenses. The limited room required to set-up a 3D printing facility and the real-time production that is made possible by this technology may overcome the need for storage and long-term stability studies of finished products, which could extemporaneously be fabricated upon request. Within hospital and compounding pharmacies, 3D printing may fully innovate galenical practice, especially impacting on clinical trials and treatment of small groups of patients (e.g. people suffering from rare diseases). Moreover, it would allow release targets (e.g. controlled release), which are not yet feasible by compounding, to successfully be addressed. In the prospect of industrial production, 3D printing would enable continuous manufacturing, of high current interest, by dramatically simplifying downstream production. Notably, 3D printing could easily be adapted not only to the manufacturing of small batches but also to that of large ones, by making more printers work in parallel or designing next-generation equipment provided with multiple printing sources.

3. 3D printing of drug products

The interest in exploitation of 3D printing within the pharmaceutical field was promoted by the development of low-cost desktop versions of printers that has made this technology widely
accessible for use in lab settings. Currently, the research activity in this specific area is mainly focused on evaluating the potential of 3D printing as an alternative approach to fabrication of drug products and as a tool for attaining innovative DDSs with unique design, composition and performance features. For example, 3D polymeric templates having a complex structure, in which a gel-based drug-containing preparation could be cast and cured, were proposed for the molding of products. Moreover, 3D printing was employed to fabricate cylindrical-shaped ABS containers, having two holes placed in different positions, to be filled with a drug formulation. By these systems, zero-order release kinetics would be attained avoiding the typical initial burst effect.

Rather than pursuing specific therapeutic goals, however, the research activity seems to be intended to highlight the versatility of the new technology, thus promoting its use for personalization of therapy.

Apart from the patent area, which is quite broad in scope with respect to 3D printing of drug products, the use of three specific techniques was actually reported in the scientific literature and, even if in rather few applications, undertaken in the industrial production: 3D printing based on solidification of powder, extrusion and, more recently, stereolithography. By way of example, a number of 3D printed drug products proposed in the scientific literature are depicted in Figure 2. These examples clearly show the degree of freedom that can be achieved in terms of shape (e.g. spherical, cylindrical, pyramidal, cubic), design (e.g. solid, hollow, pierced, polo, multilayer, coated, multi-compartment, gradient systems and relevant combinations) and composition (i.e. type of ingredients and their distribution within the product). In particular, one of more drugs can be included in whatever layer, compartment or part of the printed item. It is indeed known that, if compared with traditional manufacturing processes, 3D printing mainly offers the advantage to yield complex drug products within a single step, thus reducing the process time and the equipment needed (e.g. production and coating of the core in a single stage).
3D printing techniques currently employed for drug product manufacturing will briefly be discussed, highlighting the peculiar advantages, still open challenges and main applications proposed. A general outline of the relevant equipment is reported in Figure 3.

### 3.1 Highlights of solidification of powder

3D printing based on solidification of powder was born as an evolution of traditional 2D printing and was first developed in the early '90s at the Massachusetts Institute of Technology\textsuperscript{30-32}. It involves deposition of a liquid formulation onto a powder bed to create bonds among the particles, and a 3D solid structure is constructed by successive layering of powders. Each layer is spread over a build plate and possibly compressed by a roller (Figure 3). The binding liquid is jetted according to a 2D pattern by one or more printer heads, thermal or piezoelectric, exploiting a technology that is close to ink-jet printing. The printer head system generally moves along the horizontal plane, whereas the build plate moves down along the vertical axis, so that successive new layers of powder can be deposited. At the end of the printing process, a drying step, which is generally carried out at room temperature, and the removal of free particles are required. In addition, the product may need to be heat-treated for elimination of residual solvents and achievement of mechanical resistance. Post-process treatments, and particularly disposal of any excess of powder, require suitable facilities and equipment since the presence of fine particles might be hazardous for human health.

3D printing based on solidification of powder holds potential for the manufacturing of drug products in that it involves the use of powdered materials and binding procedures, which are largely employed in the pharmaceutical field, especially for fabrication of solid dosage forms. Moreover, as with other manufacturing techniques, it is possible to adjust the characteristics of the product by changing the process parameters. Being correlated with the thickness of powder layers, the final resolution will strictly depend on the particle size: the smaller is the latter, the more detailed will be the structure achieved. However, flowability and cohesion issues as well as adhesion of the powder particles to the equipment may have to be faced. The powder formulation could contain the API and
various excipients, including fillers, matrix formers (polymers) and release modifiers needed to define the final performance of the system.

Further critical process variables are the mode and speed of liquid deposition. This concerns the drop dimensions, movement of the nozzle and its distance from the powder bed as well as the jetting mode, which may be continuous under pressure, by drops, or alternate by means of a drop-on-demand process. The liquid can either consist in a solution or a suspension, and may contain the drug along with release-controlling polymers, one or more binders, co-solvents, plasticizers and surfactants. Not only the composition but also the rheological characteristics of the liquid should attentively be considered. Moreover, in order to selectively enhance the formation of bonds among particles in specific areas of the powder bed, the 2D wetted pattern should strictly be defined. Items are often characterized by high porosity, which could represent an advantage, such as in the case of orally-disintegrating tablets, but also an issue for the handling of products. The binder bleeding into the powder bed was identified as one of the most critical issues that may affect the product quality and performance. Indeed, bleeding along the vertical axis could cause not sufficiently resistant bonds to be created between two layers, thus reducing the mechanical strength of the final product. In addition, it may bring about poor surface quality, imprecise printing, worsened resolution and, in some cases, lack of release performance. The ability to control the moisture content of the object during fabrication is also a factor that may affect further development of this technology, and the need for post-process operations could also be seen as a limitation.

3D printing based on solidification of powder was proposed for a number of applications that are summarized in Table 2. This technique enables fabrication of systems characterized by complex geometries and different microstructure, composition as well as surface features (Figure 2). Therefore, it was particularly exploited for obtaining of pierced and multilayer products or of items provided with internal cavities, void or containing free-flowing powder formulations. The possibility of varying the spatial composition within a product, intended as the ability of introducing, with high resolution, different drugs or excipients (e.g. release modifiers, release-
controlling polymers) in predetermined inner or surface regions, was also used for one-step fabrication of DDSs enabling combined release kinetics. Analogous results were obtained by switching between binders with different composition and properties.

3D printing based on solidification of powder allows high-precision dosing that is particularly important in the case of low-dosed drugs or for the achievement of concentration gradients within a matrix system possibly leading to zero-order release\textsuperscript{39,40}. Dissolution of a drug into the binding liquid was proposed in order to increase the bioavailability of poorly water-soluble compounds, although recrystallization phenomena may occur and lead to changes in the mechanical properties and release behavior of the product\textsuperscript{33}. Besides stability issues, further critical aspects related to the drug load, limited availability of suitable binders, use of organic solvent and long process time still challenge the development of this printing technique over traditional manufacturing processes.

### 3.2 Highlights of extrusion

The 3D printing technique that exploits extrusion of molten/softened or semisolid materials as the additive process was only recently applied to fabrication of drug products, as a result of the availability of compact and relatively inexpensive equipment. Nevertheless, it has immediately shown a great potential in the field in view of well-evident similarities to pharmaceutical processing techniques such as extrusion (wet extrusion and HME) and IM. Moreover, the printing process is considered easier among customers, and can be run in do-it-yourself environments. Based on the starting materials employed, two different techniques may be identified, which make use of thermoplastic polymers, generally in the form of filaments, or of liquid/semisolid formulations, respectively.

FDM 3D printing involves the use of a thermoplastic material that is heated up and then deposited by a nozzle, in a semi-molten state, onto a build plate where it forms layers and solidifies (Figure 3). In most printers (e.g. MakerBot models), the same tractor wheel arrangement that drives polymeric filaments into the heated extrusion head also generates the extrusion pressure needed to
make the melted/softened material flow. The extrusion head is carried by a plotting system that enables 2D movements, while the build plate allows the item to grow in the third dimension. While the nozzle orifice defines the diameter of the flowing material that is deposited, the vertical lowering of the build plate imparts the nominal thickness to each layer, and they both determine the final resolution of the printed object though this is also affected by the dilatometric behavior. More recently, printers with a reverse arrangement in which the build plate moves in the horizontal plane and the extrusion head rises vertically, so that the object is fabricated bottom up, were proposed (e.g. Printrbot). After deposition of the last layer, the removal of any support material (i.e. materials used to wrap or anchor the object under construction and prevent collapses) and other post-processes (e.g. sandpapering, coating) may be needed to give the product its final aspect and properties. Some equipment are provided with multiple nozzles that allow more than one material to be processed at the same time, leading to fabrication of items having parts with different characteristics (e.g. color, composition), or the object and the support material to simultaneously be printed.

Starting materials for FDM are generally supplied in the form of filaments with defined shape and dimensions (i.e. diameter and diameter tolerances), which are produced by HME. Based on the broad demand for plastic products, filaments available on the market are mainly composed of ABS, PLA and HIPS. Exploitation of FDM in other areas, however, requires the development of new filaments composed of different materials. In this respect, not only the use of a range of polymers was explored, such as PVA, XT copolyester, nylon and TPU, but also diverse properties of filaments (e.g. color, resistance, flexibility, conductivity) were pursued. The current interest in this production technique is highlighted by the increasing number of companies focused on the manufacturing of new and high-quality filaments (e.g. TreeDFilaments) and, also, by the growing attention to safety issues, e.g. to the levels of ultrafine particles in the working area derived from the heating of PLA and ABS filaments47. However, no filaments based on pharmaceutical grade polymers are commercially available yet. In the beginning, research work was carried out using
plastics (e.g. PLA, PVA) filaments and adding APIs by soaking or re-extrusion. A few articles have recently been published on the formulation and extrusion of filaments based on pharmaceutical grade polymers such as acrylic and metacrylic acid copolymers, cellulose derivatives and EVA. Moreover, the possibility of characterizing these filaments in terms of mechanical and surface properties (e.g. stiffness, brittleness, roughness) was taken into account, using the commercially available PLA filament as a reference. Indeed, equipment used so far were conceived to work with plastic filaments available on the market. Therefore, printing issues such as break up of filaments and their wrapping around the loading gears, are likely to be encountered. Currently, the process of filament loading into the 3D printer was improved based on a trial and error approach, not only by adjusting the equipment configuration (e.g. compression force applied by the gears) but also by modifying the formulation of the filament under investigation, thus ameliorating its mechanical features. The addition of external liquid lubricants (e.g. castor oil, glycerol, oleic acid, PEG 400) to help the loading process was also described. In the future, the experience gain in the production and characterization of filaments based on pharmaceutical-grade polymers will most likely promote the development and utilization of purposely-designed printers. One of the main advantages offered by FDM in the manufacturing of drug products consists in the lack of solvents, which reduces time and costs of the process and may be beneficial to stability. The relatively high temperatures involved would limit microbial contamination. Moreover, they could promote interaction between the polymer and the drug conveyed, possibly leading to enhanced bioavailability through formation of solid dispersions or solutions. As already observed with other hot-processes, the operating conditions involved by FDM could lead, by affecting porosity, to improved release profiles in the case of prolonged-release matrices if compared with traditional manufacturing by tableting. On the other hand, as occurring in the case of HME and IM, the processing temperatures could impact on the physical stability of the object, bringing about shrinkage and warpage phenomena, and on the chemical stability of the formulation components, either excipients or active ingredients. For this reason, an accurate selection of the starting materials
is needed, and the availability of alternative polymeric filaments, which may differ in terms of physico-technological characteristics and processing conditions while allowing products with comparable performance to be obtained, could be of great interest. In addition to the physical properties of the starting materials (heat capacity, thermal conductivity, density, glass transition temperature and melt viscosity), process variables, such as loading pressure, feed rate, working temperature, deposition rate, layer height, infill percentage and number of shells, also affect the quality and resolution of the printed product\textsuperscript{22,49,56,58,60}. Moreover, the configuration of the printer (e.g., heated or unheated build plate, presence of kapton tape and nozzle orifice diameter) and the environmental conditions, particularly when the equipment is not insulated, could influence the deposition and binding of successive layers, which impacts on the mechanical resistance of the fabricated structures.

As compared with products produced by solidification of powder, objects obtained by FDM are generally characterized by greater mechanical resistance and higher degree of geometrical freedom. The interest in products endowed with such versatility in design is testified by the recent foundation of startups and academic spinouts for their commercialization (e.g. Multiply Labs, FabRx). By way of example, empty hollow structures, which can be assembled into tightly closed capsules, were produced by this technique\textsuperscript{56}. Moreover, FDM enables processing of more than one material at the same time thus allowing coated (i.e. shell-core structure) and multi-layered systems or items having a gradient composition to be fabricated, from which multiple-phase release kinetics can be achieved (Figure 2)\textsuperscript{54,58,61}.

Since a clear layer deposition pattern can frequently be identified in the finished item, the surface smoothness of printed drug products often needs to be improved, for instance by using nozzles having a smaller orifice or reducing the layer height. Resolution issues are particularly challenging when the thickness and/or presence of details in the printed item represent a critical parameter (e.g. capsular devices, coated systems)\textsuperscript{56,58}. These issues were even more evident when using equipment wherein the position of the build plate has to be manually calibrated. In this respect, new 3D
printers with automatic calibration sensors were recently developed. The release performance of the
drug product can be modulated by changing not only the filament formulation but also the density
of the deposition plot (e.g. infill percentage)\textsuperscript{48,50,57,61}. However, the main challenge of FDM is still
represented by the scaling-up and the process time, which may not be compatible with large-scale
production and advantageous with respect to more traditional manufacturing processes.

When extrusion starting from liquid and semisolid formulations is considered, Fab@Home and 3D
Bioplotter are probably the most common printers that use solutions, dispersions, gels and pastes\textsuperscript{63}
(Figure 3). In the case of 3D Bioplotter, initially developed to work in a sterile environment for the
production of scaffolds, the layers are deposited by one or more syringes under defined conditions
of temperature and pressure. On the other hand, Fab@home was the first multi-material 3D printer
available on the market via an open source. Each syringe can independently and precisely (up to 1
µL) control the deposition process through the movement of the inner plunger. One of the main
advantages related to these types of equipment is the possibility of employing a large variety of
materials, such as hydrogels, epoxy resins, chocolate and cheese, with no need for high
temperatures. However, the possible use of organic solvent could be hazardous for human health
whereas, in the best case, the use of water as the solvent involves the same critical issues described
for 3D printing based on solidification of powder (e.g. microbial contamination, drug stability,
control of moisture content during and after fabrication, drying procedures and final curing). The
resolution and mechanical characteristics of the finished item, particularly friability and hardness,
still represent a challenge for this technique. Moreover, the need for post-processes would also
increase the overall production time. The use of gels and pastes as starting materials may result in
shrinking or deformation of the printed product following the drying phase. The object may also
collapse during fabrication if layers have not sufficiently hardened to withstand the weight of the
next ones. Moreover, the technique is characterized by a low resolution as currently employed
orifices have diameter in the 0.4 - 0.8 mm range.

The main applications of 3D printing based on extrusion are summarized in Table 3 and Figure 2.
3.3 Highlights of stereolithography

As with 3D printing based on solidification of powder, which was developed starting from the ink-jet technology, SLA results from upgrade of 2D lithography to cover the third dimension\textsuperscript{73,74}. This printing process is based on solidification (curing) of successive layers of photosensitive liquid polymers (resins) through irradiation by a light source \textit{(e.g.} UV laser). Through a laser beam or a digital light projector, the specific pattern defined by the CAD file is cast over the liquid formulation (Figure 3). The light source supplies the energy required for promoting cross-linking of the polymeric chains, thus leading to the formation of solid layers with a defined resistance, the first one adhering to the build plate. Subsequently, the plate is lowered, the first layer is coated with another fraction of liquid resin, and the curing step is repeated. Therefore, printed items are generally built bottom-up from a build plate that is positioned just below the surface of the liquid. However, a top-down approach was recently proposed, wherein the light is projected from underneath, through a transparent, non-adhering plate. In this case, the build plate is dipped in the liquid resin from above. Although up-to-bottom built structures are subject to higher stresses in that they need to be separated from the bottom build plate after each deposition step, this approach enables the use of smaller amounts of resin, better protection of the object surface from the environment and attainment of a smoother surface.

In view of its suitability for fabrication of precise geometries and internal architectures with details in the micrometer range, SLA is particularly useful in the manufacturing of scaffolds. Indeed, it yields printed objects having up to 20 µm resolution, as compared with the 50-200 µm range granted by other additive manufacturing techniques, along with high precision and smooth surfaces. However, post-curing with UV light is often required to improve the mechanical properties of the printed objects. The thickness of deposited layers is determined by the viscosity and spreading of the liquid resin employed and the energy of the light, which is mainly controlled by the power and scanning speed for laser systems and by the exposure time for projection systems, respectively.
The main challenge for SLA technique is related to the limited availability of suitable photopolymers and their quite complex kinetics of curing reactions. Very few materials have been developed so far, and most of them are expensive, toxic, smelly and need to be shielded from light to avoid premature polymerization. Moreover, none is categorized as GRAS and could safely be used for the manufacturing of drug products. Only recently, some biodegradable resins have been identified (e.g. PCL, chitosan) and proposed for fabrication of scaffolds\textsuperscript{75,76}. However, since the temperature increase during printing is limited, this technique has a great potential for processing of thermally labile drugs. The few applications of 3D printing based on SLA are summarized in Table 4.

4. Regulatory perspectives

Beyond the identification of major targets and development of appropriate techniques for fabrication of drug products, the main challenge for personalized pharmacological therapy would most likely be related to the regulatory issues involved\textsuperscript{78}. These may primarily concern health hazards and safety aspects resulting, for instance, from the use of fine powders, organic solvents and irradiation, as well as from possible formation of unknown degradation products derived from high-temperature processing even of already approved materials. Approval of a sufficiently broad variety of thermoplastic polymers and liquid resins to be used in FDM and SLA, respectively, also constitutes a fundamental step that may strongly limit profitable exploitation of 3D printing in the manufacturing of medicines. Moreover, the actual possibility of undertaking personalized therapy would necessarily depend upon implementation of production models that may enable to efficiently turn the therapeutic needs of individual patients into small batches of appropriate drug products. It is well known that the FDA regulatory approach is currently based on mass production, standardization and batch validation. So, the vast majority of rules applied so far would hold poor promise for the manufacturing of customized medicines. In this respect, FDA has started to consider the impact of 3D printing on the production of medical devices. It is established that all of them,
including those produced by additive manufacturing techniques, have to meet the same quality and compliance requirements \(i.e.\) Quality System. Since regulation addresses many different types of devices, it cannot provide specific prescriptions for each fabrication process and rather sets out a framework to ensure that these products consistently meet applicable specifications. Therefore, the trend is to include 3D printed devices under the existing regulations while evaluating similarities and differences over traditionally manufactured products and identifying, through a joint work with manufacturers, the most critical points that need to strictly be controlled. Such a policy led to the drafting of shared Technical Considerations for Additive Manufactured Devices that have recently been published\(^7\).

With respect to 3D printed drug products, no tangible progress has been made so far, although FDA is presently engaged in a workable regulatory platform that promotes innovation while ensuring accuracy by issuing discussion papers, holding workshops and collaborating with stakeholders. Indeed, the exploitation of 3D printing techniques for the manufacturing of drug products represents a very challenging subject. For instance, depending on the site where the printing process is run \(e.g.\) compounding pharmacy, hospital pharmacy, pharmaceutical industry), the resulting product should be classified among individual extemporaneous, hospital or industrial preparations, and each of them might have to follow specific regulatory paths. On the other hand, independent of the fabrication site, their quality must be ensured. Therefore, all aspect entailed by the manufacturing process has to be comprehensively considered such as hardware, software, raw materials \(e.g.\) filaments) and their source, operators and training, responsibilities, outcomes and quality controls. Moreover, when the 3D printing process is carried out at the point of care, high-quality software and hardware supplied by specialized companies \(e.g.\) printers allocated within a high-controlled production chamber), specifically developed for pharmaceutical use, would be needed.

In the meantime, the first 3D printed tablet, Spritam\(^\text{R}\), fabricated by powder deposition-based Aprecia Zipdose\(^\text{R}\) technology, was approved in summer 2015. This product is available in different strengths up to 1000 mg and enables a new mode of administering anti-epilepsy drug levetiracetam.
Its goal, however, is only marginally connected with personalization. In this case, 3D printing was indeed exploited for the purpose of fabricating a porous dosage form with high drug load, able to fast disintegrate in a small amount of liquid, which could be critical to attain by traditional manufacturing techniques. Hence, it deals with a dosage form produced on a large-scale and approved via the regulatory procedure currently in use. A recent study also demonstrated bioequivalence of this product in the fasted state to the conventionally fabricated immediate-release tablets available on the market\textsuperscript{80}. 
References


52. Holländer J, Genina N, Jukarainen H, Khajeheian M, Rosling A, Mäkilä E, Sandler N  
2016. Three-dimensional printed PCL-based implantable prototypes of medical devices for 

acetate (EVA) as a new drug carrier for 3D printed medical drug delivery devices. Eur J 
Pharm Sci 90:53-63.

extruded filaments based on pharma-grade polymers for 3D printing by Fused Deposition 

55. Pietrzak K, Isreb A, Alhnan MA 2015. A flexible-dose dispenser for immediate and 

fused deposition modeling (FDM) of a swellable/erodible capsular device for oral pulsatile 

Adaptation of pharmaceutical excipients to FDM 3D printing for the fabrication of patient-

a shell-core delayed release tablet using dual FDM 3D printing for patient-centred therapy. 

improve the printability of and regulate drug release from pharmaceutical solid dispersions 
prepared via fused deposition modeling (FDM) 3D printing. Eur J Pharm Biopharm  
108:111-125.

60. Aho J, Boetker JP, Baldursdottir S, Rantanen J 2015. Rheology as a tool for evaluation of 


Table 1: classification of the main additive manufacturing techniques

<table>
<thead>
<tr>
<th>TECHNIQUE</th>
<th>SUBSTRATE</th>
<th>MECHANISM OF LAYERING</th>
<th>ADDITIVE PROCESS</th>
</tr>
</thead>
<tbody>
<tr>
<td>FUSED DEPOSITION MODELING</td>
<td>filament (thermoplastic polymer)</td>
<td>melting/softening by an heated nozzle</td>
<td>extrusion</td>
</tr>
<tr>
<td>ELECTRON BEAM DIRECT MANUFACTURING</td>
<td>wire (metal)</td>
<td>melting by an electron beam</td>
<td>direct energy deposition</td>
</tr>
<tr>
<td>SELECTIVE LASER SINTERING</td>
<td>solid particles</td>
<td>melting by laser</td>
<td>solidification of powder</td>
</tr>
<tr>
<td>THREE DIMENSIONAL PRINTING</td>
<td></td>
<td>binding by wetting</td>
<td></td>
</tr>
<tr>
<td>STEREOLITHOGRAPHY</td>
<td>liquid (photopolymer)</td>
<td>binding by UV ray</td>
<td>photopolymerization</td>
</tr>
<tr>
<td>LAMINATED OBJECT MANUFACTURING</td>
<td>sheets (paper, metal, plastics)</td>
<td>cutting by laser</td>
<td>sheet lamination</td>
</tr>
</tbody>
</table>
Table 2: overview of dosage forms fabricated by 3D printing based on solidification of powder

<table>
<thead>
<tr>
<th>Product</th>
<th>Performance</th>
<th>Starting materials</th>
<th>Equipment</th>
<th>References</th>
</tr>
</thead>
</table>
| Implantable systems/inserts   | Immediate, prolonged, pulsatile release and combinations of differing kinetics | **POWDER**
PCL, PLA, PLGA<br><br>**BINDING LIQUID**
SOLVENT: acetone, ethanol, methanol, water<br>API: ethinyl estradiol; isoniazid; levofloxacin; rifampicine | Custom-made (Fochif Mechatronics Technologies Co. Ltd, Therics)                          | 37, 41-44  |
| Oral systems                  | Immediate and fast release                               | **POWDER**
colloidal silicon dioxide, HPMC, Kollidon® SR, lactose, MCC, PVP, SLS, stearic acid<br>API: acetaminophen<br><br>**BINDING LIQUID**
SOLVENT: acetone, ethanol, water<br>BINDER: EC, Eudragit® E, RL and RS, PVP, TEC, Tween 20<br>API: acetaminophen, diclofenac, chlorpheniramin maleate, fluorescein disodium salt (tracer) | Custom-made (Fochif Mechatronics Technologies Co. Ltd, TheriForm™)                      | 36, 39, 40, 45, 46 |
| Orally disintegrating tablets | Immediate and fast release                               | **POWDER**
colloidal silicon dioxide, crosslinked PVP, lactose, maltitol, maltodextrin, mannitol, PVP, API: acetaminophen<br><br>**BINDING LIQUID**
SOLVENT: ethanol, phosphate buffer (pH = 4), water<br>BINDER: ascorbic acid, citric acid, EDTA, PVP, Tween 20<br>API: alizarin yellow (tracer), captopril, methylene blue (tracer) | Custom-made (Fochif Mechatronics Technologies Co. Ltd)                                  | 33-35       |
Table 3: overview of dosage forms fabricated by 3D printing based on extrusion

<table>
<thead>
<tr>
<th>Product</th>
<th>Performance</th>
<th>Starting materials</th>
<th>Equipment</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>EXTRUSION OF SOFTENED/MOLTEN MATERIAL</td>
<td></td>
<td></td>
<td></td>
<td>52, 53, 64, 65</td>
</tr>
<tr>
<td>Implantable systems/Inserts</td>
<td>Prolonged release</td>
<td>EVA, PCL, PLA, PLGA, PCL, API: 5-fluorouracil, gentamicin sulfate, indomethacin, methotrexate</td>
<td>Commercial (MakerBot, Prusa Research)</td>
<td>66</td>
</tr>
<tr>
<td>Dermal patches</td>
<td></td>
<td>PCL, commercial PLA filament (Flex EcoPLA™), commercial TPU filament (NinjaFlex™), API: salicylic acid</td>
<td></td>
<td>48, 51, 55, 61, 66-69, 26, 57, 50, 58</td>
</tr>
<tr>
<td>Oral systems</td>
<td>Immediate, prolonged, pulsatile, delayed release and combinations of differing kinetics</td>
<td>commercial PVA filament, crospovidone, EC, Eudragit® E, L, RL and RS, low viscosity HPC, HP, HPM, lactose, MCC, PVP, Soluplus®, TEC, triacitin, tribasic phosphate, tricalcium phosphate, talc, API: 4-aminosalicylic acid, 5-aminosalicylic acid, acetaminophen, budesonide, caffeine, captopril, diclofenac sodium, dipyridamole, fluorescein (tracer), prednisolone, theophylline</td>
<td>Commercial (Fab@Home, (RegenHU)</td>
<td>70-72</td>
</tr>
<tr>
<td>Capsule shells</td>
<td>Pulsatile release</td>
<td>HPC, PEG</td>
<td></td>
<td>56</td>
</tr>
<tr>
<td>EXTRUSION OF LIQUID/SEMISOLID MATERIAL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oral systems</td>
<td>Immediate, prolonged release and combinations of differing kinetics</td>
<td>croscarmellose sodium, HPMC, lactose, mannitol, MCC, PVP, sodium starch glycolate, PEG, tromethamine, SOLVENT: acetone, dimethyl sulfoxide, ethanol, water, API: acetylsalicylic acid, atenolol, captopril, glipizide, guaifenesine, hydrochlorothiazide, nifedipine, pravastatin, ramipril</td>
<td>Commercial (Fab@Home, (RegenHU)</td>
<td></td>
</tr>
</tbody>
</table>
Table 4: overview of dosage forms fabricated by 3D printing based on stereolithography

<table>
<thead>
<tr>
<th>Product</th>
<th>Performance</th>
<th>Starting materials</th>
<th>Equipment</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oral systems</td>
<td>Prolonged</td>
<td>DPPO, PEG, PEGDA</td>
<td>Commercial (Formlabs Inc.)</td>
<td>77</td>
</tr>
<tr>
<td></td>
<td>release</td>
<td>API: 4-aminosalicylic acid, acetaminophen</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dermal patches</td>
<td>Prolonged</td>
<td>DPPO, PEG, PEGDA</td>
<td></td>
<td>66</td>
</tr>
<tr>
<td></td>
<td>release</td>
<td>API: salicylic acid</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Figure 1:** outline of common steps generally entailed by a 3D printing process

**Figure 2:** outline of 3D-printed drug products proposed in the scientific literature (references in brackets)

**Figure 3:** outline of equipment relevant to 3D printing based on powder deposition, extrusion of molten/softened or semisolid materials and stereolithography
Figure 1: outline of common steps generally entailed by a 3D printing process

69x25mm (600 x 600 DPI)
Figure 2: outline of 3D-printed drug products proposed in the scientific literature (references in brackets)

129x88mm (600 x 600 DPI)
Figure 3: outline of equipment relevant to 3D printing based on powder deposition, extrusion of molten/softened or semisolid materials and stereolithography

68x24mm (600 x 600 DPI)