Microneedles: An advancement to transdermal drug delivery system approach

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ABSTRACT
Novel drug delivery system offers several advantages which could outweigh the benefits of other drug delivery systems. The transdermal drug delivery system being one of them offers supremacy by by-passing the first pass metabolism which eventually helps in eradication of gastrointestinal irritation. However, the major drawback of the transdermal drug delivery system is the hindrance created via the stratum corneum. This protective barrier of the skin does not allow required penetration of the drug via skin into the systemic circulation. Thus, in order to overcome this hurdle, a replacement to this type of novel drug delivery system, namely, “microneedle drug delivery system” helped to improve various pitfalls of transdermal drug delivery system, such as skin barrier function, restrictions toward using of specific drugs only, bioavailability, patient compliance, diffusion rate, and plasma concentration level. A microneedle drug delivery system, thus, is advancement to transdermal drug delivery system which includes delivery of drug via microneedle into systemic circulation, thus increasing patient compliance and avoiding problems rendered by transdermal drug delivery system.

INTRODUCTION
Transdermal drug delivery system includes the delivery of drug via superficial lipophilic layer of the skin stratum corneum which is 10–15 µm thick. Stratum corneum obstructs the entry of antigens, bacteria; thus defining the skin barrier function which eventually suppresses the percutaneous penetration. Henceforth, the utmost challenge included a search for an alternative way which would surpass the repercussions provided by the skin barrier function of stratum corneum (Bhowmik et al., 2013). Various innovations, such as iontophoresis, electroporation, sonophoresis, magnetophoresis, and use of penetration enhancers have been proposed by different researchers which could help elevate the permeation and delivery of drug (Bhowmik et al., 2013). Out of all, the concept of using “microneedles”—micron-sized needles for the delivery and penetration of the drug is considered to be the most efficient method. The proof to this has been the experiments performed and subsequent results obtained by researchers and scientists (Bhowmik et al., 2013). A microneedle does not limit itself by improving only drug penetration. It has several other significant positive aspects (Wang et al., 2017). The drug delivery is specially preferred for pediatric application (Duarah et al., 2019) Thus, this review article focuses on microneedle drug delivery—its existence by overcoming the short comers of transdermal drug delivery system, types of microneedles used, different approaches for delivery, fabrication of microneedles, mechanism, materials, evaluation, and specific applications of microneedle drug delivery system.

Conventional mode of drug delivery
The delivery of the drugs plays a vital part in medicine and paramedical field. Normally, drug delivery system can be referred to as various ways, methods, and approaches for delivering the particular drug at proper site of action in order to obtain the highest amount of desired therapeutic effect. Normally, conventional modes of delivery include oral route, usage of carrier injections, or using novel approaches like transdermal drug delivery.
system for delivery of drug. With growing need for painless and less infectious prone way of drug delivery, microneedles are prone to fit the void for satisfying the requirement (Kaushik et al., 2001; Kshirsagar, 2000).

Transdermal drug delivery system

Transdermal drug delivery system (TDDS) is described as a novel drug delivery approach in which discrete and appropriate quantity of drug is delivered through the topmost layer of the skin which releases drug at a controlled rate for prolonged period of time into the systemic circulation. It can be described as a type of novel drug delivery approach which follows zero order drug release pattern (Kshirsagar, 2000).

The types of transdermal drug delivery include -

Membrane permeation controlled TDD system

In this type of transdermal drug delivery system, drug reservoir is in between backing layer and polymeric membrane. As shown in Figure 1(a), the drug reservoir area consists of the drug which is dispersed in the polymeric matrix solution to form a suspension like paste which is then delivered at a controlled rate through the rate controlling membrane which is non-porous or micro porous (Patel et al., 2012). Drugs like estradiol are delivered through membrane penetration (Chetkowski et al., 1986).

Matrix diffusion controlled transdermal drug delivery system

As shown in Figure 1(b), occlusive base plate, drug reservoir, polymeric membrane is present. The drug reservoir consists of the drug which is surrounded/coated with the matrix. Here, matrix could be any water-loving (hydrophilic) molecules or even lipophilic molecules. The dispersed form of drug and matrix forms a medicated disc like structure from which required amount of drug gets released at a controlled rate into the systemic circulation. The drug is in solution form in matrix and is backed by adhesive layer (Banerjee et al., 2014). Drugs like nitroglycerine can be delivered using this pathway (Rayment et al., 1985).

Reservoir gradient controlled transdermal drug delivery system

This type helps in dealing the issues related with non-zero release profile of the drug (Kesarwani et al., 2013). In this the drug and polymer matrix combined together in a reservoir could be converted in the form of a reservoir gradient along the diffusional pathway across the layer which leads to delivering a drug at a controlled rate as described in Figure 1(c). Glyceryl trinitrate is the drug which forms a depot system for controlled delivery (Patel et al., 2012; Wiedersberg et al., 2014).

Micro reservoir dissolution controlled transdermal drug delivery system

This type of drug delivery system is depicted in Figure 1(d) which has the combination of matrix and reservoir dispersion type of drug delivery system which includes two basic steps—Forming the drug reservoir which is done by suspending the drug in water soluble or aqueous solutions and then mixing/ dispersing the suspension with lipophilic polymer by strong mechanical force which results in the formation of numerous microscopic reservoirs and finally the release of the drug at controlled rate (Patel et al., 2012).

Transdermal patches are widely used. Several transdermal patches available are for Pain relief, Patches for reducing hypertension, Vitamin B12 patches, Nicotine patches, etc. Trade name of certain transdermal products include Catapres-TTS®, Climara®, Climara pro®, CombiPatch®, Duragesic®, Menostar®, Ortho Evra®, Oxytrol®, Transderm scop®, Vivelle®, Vivelle-Dot® (Wokovich et al., 2006). These transdermal products provide several benefits like it provides prolong effect with possibility of self-administration (Chaudhary et al., 2013), provides an alternative for patients who are unable to take oral medications, elimination of first past hepatic metabolism, avoid problems faced due to parenteral therapies, requires less dosage frequency (Allen and Ansel, 2013), visual confirmation while patch...
is applied, useful for trypanophobes (Raza et al., 2015), avoidance of enzymatic degradation, and gastrointestinal irritation. Also, it can be an alternative for the patients who are unable to take oral medications. Another advantage is that the drug release stops as soon as patch is removed. Despite of these ascendencies, there occur several stumbling blocks like unsuitableness for the drugs that irritate or sensitize skin and allowance for relative potent drugs only due to limitations related with skin permeability, problems related with non-adherence of a patch to skin, uneasiness while wearing a patch, change in barrier function of skin from site to site and person to person, the thickness of stratum corneum, poor solubility, allowance for only non-ionic, lipid and water-soluble molecules of particular molecular size, and shape limits the use of this type of drug delivery system. Hence, the more innovated type of drug delivery like microneedle drug delivery system can be employed to replace the setbacks offered by TDDS. Also, the transdermal drug delivery is unable to pass through stratum corneum which is the biggest limiting factor (Lampe et al., 1983; Ledger, 1992). The different transdermal systems currently available in the market, which could also be delivery with more effectiveness by microneedle drug delivery systems, are mentioned in Table 1.

Microneedle drug delivery system

The overarching response to microneedles is due to the concept of disrupting the stratum corneum layer of skin which can lead to smooth pathway for the entry of desired molecules. A microneedle drug delivery system can, thus, be explained as a Novel drug delivery system approach in which drug is delivered into the systemic circulation through the needles. The system in which micron-sized needles pierce the superficial layer of the skin and diffuses the drug across the epidermis layer which is then passed on into the blood capillary region for active absorption (Silpi et al., 2011) as microneedles are short and thin they help in avoiding the pain caused to patient (Shilpa et al., 2001). This could be achieved by Microfabrication technology (Blushan and Caspers, 2017; Donnelly et al., 2010). With passing time, different materials, such as dextrin, stainless steel, ceramic, and maltose were utilized for preparing the microneedles (Donnelly et al., 2010). Few years later, pattern of permeability of drug to the skin via microneedles was studied which lead to a conclusion that permeability factor through microneedles could be elevated to three to four times as compared to that of TDDS. It was observed that the skin barrier function could be overcome by using micron-sized needles. Later, different materials through which microneedles could be fabricated were introduced. Hence, this lead to the inception of microneedles as a solution for more productive results in terms of bioavailability and permeation of drug. Scientists put down efforts using technology for proper optimization and geometrical measurements required for achieving accurate insertion in the skin of human which was major goal with respect to microneedle (Kong et al., 2011).

TYPES OF MICRONEEDLES

Solid microneedle

These microneedles use the passive diffusion path (Larraneta et al., 2016). Solid microneedles follow two steps, as shown in Figure 2, in which the microchannels in the skin are first created by inserting the microneedle, followed by allowing the drug formulation to pass over the generated microchannels (Kaur et al., 2014). Solid silicon microneedles proved to be the most prominent approach because of their sufficient biocompatibility (Wei et al., 2010). The manufacture of such a type of microneedle involves the use of various materials, such as polymers (Park et al., 2006), stainless steel metal (Martanto et al., 2004), biodegradable polymers (Park et al., 2005), and silicon (Li et al., 2010), for which silicon is the most important material to be used. Food and Drug Administration must approve these materials before fabrication starts on the basis of biocompatibility and safety (Kotzar et al., 2002). For proper production, various methods are used. One possibility is the use of thermal oxidation and lithography, wherein silicon on insulator wafer and oxide layer is processed to form a silicon oxide layer by thermal oxidation and lithography

![Figure 2. Mechanism of delivery of drugs by four types of Microneedles solid, coated, dissolving and hollow.](image)

<table>
<thead>
<tr>
<th>Drug Used</th>
<th>Type of disease treated</th>
<th>Type of TDD used</th>
<th>Advantages</th>
<th>Side effects</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Estradiol</td>
<td>Postmenstrual syndrome</td>
<td>Membrane</td>
<td>Reduce problems related with menopause</td>
<td>Breast tenderness</td>
<td>(Utian, 1987)</td>
</tr>
<tr>
<td>Scopolamine</td>
<td>Motion sickness</td>
<td>Matrix</td>
<td>Reduces motion sickness</td>
<td>Oral administration</td>
<td>(Clissold and Heel, 1985)</td>
</tr>
<tr>
<td>Testosterone</td>
<td>Hypogonadism</td>
<td>Reservoir</td>
<td>Increase muscle mass</td>
<td>Sleep apnea, skin reaction</td>
<td>(Wang et al., 2000)</td>
</tr>
<tr>
<td>Rivastigmine</td>
<td>Dementia</td>
<td>Microreservoir</td>
<td>Alzheimer’s Disease</td>
<td>Spinning sensation</td>
<td>(Kurz et al., 2009)</td>
</tr>
<tr>
<td>Fentanyl</td>
<td>Pain relief</td>
<td>Reservoir</td>
<td>Decrease in oral mucositis pain</td>
<td>Convulsions, dry mouth</td>
<td>(Kim et al., 2005)</td>
</tr>
</tbody>
</table>
to obtain the first layer. The second layer (parylene coated) is deposited thereon to protect and isolate the chip. Furthermore, the microneedle electrode assembly is emancipated by separating the silicon oxide layer. Thus, the silicon array is obtained by modifying the parylene layer, which was previously under the silicon oxide layer, as a carrier layer of the solid silicon microneedle obtained by simply depositing the parylene layer to impart adhesives and strength to the microneedle (Wang et al., 2012). The main precision that emanates from a solid silicon microneedle is the rare occurrence of inflammatory or infectious conditions (Vicente-Perez et al., 2017). However, the high silicon cost is causing a setback in the use of silicon as the material for solid microneedles (Yue et al., 2009). Thus, a solid microneedle can also use other materials, such as parylene (Johnson and Wise, 2010), polyimide (Seo et al., 2004), nickel (Wang et al., 2007), titanium (Fofonoff et al., 2002), Polyethylene methacrylate, and polyvinyl (Matteucci et al., 2009).

Drug-coated microneedles

These type include coating of the drug on the solid microneedle before inserting it into the skin (Gill and Prausnitz, 2007a). The coated microneedle is inserted into the skin where the drug gets dispersed into the systemic circulation. This system is utilized to deliver complex and macro molecules, including Deoxyribonucleic acid, drugs, like desmopressin, parathyroid hormone, and vaccines (Gill and Prausnitz, 2007b, Quinn et al., 2014), peptides (Cornier et al., 2004), proteins (Matriano et al., 2002). Different methods like micro dip-coating for vaccine delivery (Koutsonanos et al., 2009) are used for fabricating microneedles onto which a drug solution can be coated to obtain a drug-coated microneedle ready for administration. Other method for developing coating microneedle include fabrication via laser cutting in which stainless steel material is used and according to the required acreage the sheet is cut by allowing infrared rays to pass through it. The obtained cutout is then cleaned by using proper cleanser. Furthermore, the electro polishing is required which is done by using copper plate which acts as a cathode and microneedles to be obtained would act as anode. The procedure requires continuous vibration in order to avoid the unwanted air bubbles. Microneedles are then cleansed using the de-ionized water followed by hot water which are then allowed to dry and the stored at optimum temperature. A coating solution is then prepared of the drug to be coated and microneedle is then coated using dip-coating technique. Coated microneedle is specifically used in treating enuresis, diabetes insipidus, hemophilia A, Willebrand’s disease and certain trauma induced diseases (Gill and Prausnitz, 2007a).

Dissolving microneedles

These type of microneedles include usage of the biocompatible polymers like poly vinyl alcohol (Chu et al., 2010), carboxymethyl cellulose (Guo et al., 2013), or sugars (Miyano et al., 2005) which on insertion into the skin releases the drug and dissolves into the systemic circulation due to which a controlled release is obtained. The needle or the coating on the needle dissolves itself to release drug (Lee et al., 2008; 2011; Matriano et al., 2002). The main advantage is low cost of polymers and usage of water-soluble materials. But, one of the major downsides is deposition of polymers into the skin. These are prepared by lithographic approach (Moga et al., 2013) or a two-layered approach in which two plates are used. Drug and polymer are mixed and the solution formed is allowed to pass into the holes of the array. The push and pull movement is enforced onto the two plates placed parallel to each other. The polymeric solution is allowed to pass into plate below and the top plate is removed at the end after the dissolving microneedle have been fabricated as the polymeric solution gets solidified resulting in conical shaped microneedles which can be mostly used in administering influenza vaccine (Roxhed and Griss, 2008; Sullivan et al., 2008). Recently, Wu et al. (2016) prepared sinomenin loaded dissolving microneedles using a specialized fabrication technique using biocompatible polymers. (Wu et al., 2008).

Hollow microneedles

These microneedles are the one which allows the passage of the drug through the hollow bore channel present in the needle. The needle is inserted into the skin and the drug is allowed to pass through the bore which diffuses into the systemic circulation (Kurz et al., 2009). It is made up of various materials, such as silicon, metals (Chandrasekaran and Frazier, 2003; Roxhed and Griss, 2008; Roxhed et al., 2008), glass (Wang et al., 2006), and polymers (Samoura et al., 2007). The main obstacle of hollow microneedle is the problem of clogging of the bore which leads to resistance to flow (Gardeniers et al., 2003). It can be manufactured by using lithographic technique which ultimately helps in intradermal drug delivery (Laurent et al., 2007). For fabrication, a silicon wafer is used onto which the photoresist layer is spun and the deep molds are formed after the ultraviolet rays are allowed to pass through the molds. By wet etching process, the KOH solution is used to which silicon wafer gets etched and the required microneedles are obtained by cutting silicon wafer into a microneedle array (Torin et al., 2011). Hollow microneedles are utilized in extraction of the blood sample by least amount of invasion. It helps to replace traditional technique of utilizing hypodermic needle (Li et al., 2013). The mechanism of various microneedles is described in Figure 2.

Approaches

Each and every microneedle whether it is solid or drug coated follow certain common steps in mechanism. Microneedles when inserted leads to mechanical disruption of the skin creating pore, and then, the drug coated on the needle is dispersed in the systemic circulation. Microneedles firstly disrupt the stratum corneum region of the skin. Stratum corneum includes skin barrier function (Bouwstra and Ponec, 2006) and microneedle helps to eradicate the problem of skin barrier function of stratum corneum. Hence, microneedles create a pathway for drug to directly get delivered into the blood rich region bypassing the stratum corneum which is not the case in transdermal drug delivery system. To give the desired outcome, it follows poke and patch, coat and poke, poke and release, poke and flow approach as discussed in Table 2. It depends on the fabrication of a microneedle which can be in-plane or out-plane. In-plane microneedles are parallel to the surface and out-plane microneedles are perpendicular to the surface as shown in Figure 3 (Van der Maaden et al., 2012).
Clinical Trials on Microneedle Patch

In order to understand the working and advantages of microneedle patches over the hypodermic needle or any other drug delivery system like transdermal drug delivery, Arya et al. (2017) worked for many years to develop a technology based on microneedle and finally designed a technique for delivery by recommending use of the microneedle patch. Arya et al. (2017) worked at Georgia Tech and Emory University who designed microneedle patch and manufactured them at Global Centre for Medical Innovation at Atlanta. Individuals between the age group of 18 to 57 were chosen as the participants for the clinical trials. Seven males and seven females were considered for each category of the test. The test was divided into several parts (Arya et al., 2017).

The effectiveness of microneedle patches: The study was conducted on four groups including the people with traditional flu shot as first group, microneedle patch (vaccine) as second group, placebo patch without drug as third group, and people who administered the patch using the instructional video as fourth group. The results obtained were quiet interesting as it was observed that the patch worked similar to the traditional flu shot. The drug bioavailability was monitored and it was same to that of first group. Hence, the patches were approved for further studies (Arya et al., 2017).

Skin toxicity studies: The skin scoring scale was utilized in which the site at which the patches applied was evaluated on the factors, such as pain, tenderness, erythema, and swelling. Pain was evaluated on the basis of after effects, i.e., the effect after the patch is removed. Similarly, tenderness was defined on the basis of touch on the skin by the evaluator after the patch is administered and the appropriate results were obtained. (Arya et al., 2017).

For the evaluation through skin staining and microscopy, 1% gentian violet solution was used to stain the skin. After the insertion of microneedle, it was immediately removed off and the amount of stained skin was observed with the help of microscope and the punctures that microneedle made was counted, i.e., the stained part. Skin staining is calculating the microneedles that penetrated into the skin. The efficiency of microneedles was checked, i.e., microneedles that actually delivered the drug and dissolved after administration. The results obtained were satisfying as it was observed that almost 95% of the microneedles penetrated in the skin, i.e., punctured the skin layer and delivered almost 70%–80% of the drug after penetration (Arya et al., 2017).

To evaluate the microneedle patches based on user surveys, participants had to answer short questions based on patches. The pain during administration, the confidence, and preference of a conventional patch or a microneedle patch were evaluated. As a result, the evaluators found that 86% were confident in applying the patch and that the majority of participants experienced no pain. In addition, it was observed that the participants preferred more microneedle patches compared to conventional injections because the patches did not cause pain and were easy to handle and use (Arya et al., 2017).

The advancement in microneedle drug delivery can be judged from the success of these microneedles in delivering drug into systemic circulation. Few of these successful systems are listed in Table 3.

Applications of Microneedle Drug Delivery System

The some of the important applications of microneedle drug delivery along with the successful product are listed in Table 4.

Microneedles for vaccine delivery

The increase in awareness among people regarding vaccination has led scientists to find new approaches for the delivery of vaccine. The recent one being microneedle drug delivery in which the antigen is directly introduced in the dermis. Thus, immune response can be achieved and vaccination therapy can be completed. This activity is carried out mostly by using dissolving type of microneedles. Various techniques and devices like “mantoux” or “soluvia microinjection system” can be used to perform this process. According to the studies, mantoux tecique is difficult to perform as it requires professional person, whereas soluvia technique is commonly used and that too for the delivery of influenza vaccines, such as trivalent or quadrivalent or influenza type A or B (Pettis and Harvey, 2012; Van Damme et al., 2009; Vankerckhoven and Van, 2010).
Microneedles for insulin delivery

Many research studies were carried out for insulin delivery by various scientists, such as Martanto et al. (2004) and Henry et al. (1998), and they found that the insulin delivered through microneedle drug delivery was appropriate and it produced proper biological effect and maintained the blood glucose level. Also, bioavailability of insulin was found to be proper as insulin was introduced into the body by coating it on the microneedle which pierced the stratum corneum layer which led to the delivery of insulin. The first approach for insulin delivery was done by using 10-IU standardized insulin lispro which showed good absorption rate (Pettis and Harvey, 2012). Borosilicate glass microneedles were used (Gupta et al., 2009; 2011).

Microneedles for delivery of parathyroid hormone

It is used for the delivery of the parathyroid hormone, i.e., teriparatide. Drug-coated type of microneedles is used in it which delivers the drug after it is inserted in the skin, i.e., the drug coated gets dissolved after insertion. The delivery of parathyroid hormone through microneedle is not yet approved. But, it has undergone phase 1 and 2 studies which concluded that the drug delivered via microneedles reached three times earlier to the maximum concentration as compared to that delivered via traditional route (Pettis and Harvey, 2012). Parathyroid patches have also been prepared (Daddona et al., 2011).

Microneedles for delivery of Lidocaine and other anesthetics

The type of microneedles used here were solid and hollow microneedles. The anesthetic drug was delivered to the various participants selected for study and criteria, i.e., pain and numbness obtained were evaluated. The results showed that the pain was less as compared to that of traditional needle and numbness obtained was as effective as that of the conventional way of approaching numbness to particular part. Thus, it was concluded that microneedles were more effective than hypodermic needle in producing more numbness and less of pain which consequently worked for the betterment of patient (Pettis and Harvey, 2012). Solid-based microneedle has been formulated for delivery of lidocaine for patients to be anesthetized (Zhang et al., 2012).

Microneedles for cosmetic purpose (Dermaabrasion)

Microneedles are used for decreasing the scars, wrinkles, etc from the skin. This is achieved by puncturing the skin multiple times through microneedles, earlier which was done through hypodermic needle. The pores, thus, created through microneedles resulted in the increase of collagen growth and breaking or disruption of old collagen which caused damaged to the skin. This led to decreasing the wrinkles, also helped in treating acne. The special instrument made for this is Dermaroller®. It is 12 cm long and consists of 192 miniature needles arranged in 24 circular arrays. The length of which is 0.5 to 3 mm and diameter of which is 0.1 to 0.25 mm. These needles can be adjusted according to the depth, skin layer in which one wants the insertion (Fabbrocini et al., 2009; Nair and Arora, 2014; Pettis and Harvey, 2012).

Microneedles for naltrexone

Microneedle patches were used to treat opioid-dependent patients as well as alcohol addict treatment. When patches were administered, it was found that the optimum plasma level was obtained within 2 hours after administration and the effect prolonged till 72 hours. Thus microneedles coated with naltrexone, thus delivered drug into the systemic circulation which helped block the opioid effect (Pettis and Harvey, 2012; Stinchcomb et al., 2008).

Microneedles for acne scar treatment (Dermaroller)

For acne scar treatment, microneedles are used. The area with acne scar is treated with anesthetic agent and the Dermaroller is used and allowed to move in vertical and horizontal direction. The saline pads are used for treating the bleeding which could be controlled. The process takes 20 minutes for getting completed. Also, home care derma rollers are used for delivering anti-ageing products (Dogra et al., 2014).

Microneedles for glucose monitoring

Microneedles are used indirectly for glucose monitoring. Previously, devices, namely, “Cygnus glucowatch” were used more for monitoring the glucose levels in the body (Tierney et al., 2001). Using silicon microneedles gave proper precision, appropriate needle penetration which helped extracting interstitial fluid with minimum pain which helped monitor blood glucose level. (Koschinski and Heinemann, 2001) Commercially available glucose monitoring/sensing device is Kumetrix® which is made up of silicon microneedle (Smart and Subramanian, 2000).

Microneedle patch for iron deficiency anemia

Success in modulating microneedle patches to replace the oldest approach of using iron supplement and parenteral approaches for treating iron deficit human was achieved by using micro molding technique which helped treat anemia in patients without any gastric side effect. Rapidly dissolving type of microneedles loaded with ferric pyrophosphate which could help treat Iron deficiency anemia efficiently was formulated. On carrying out in vivo and in vitro studies, satisfying results were obtained (Maurya et al., 2018).

Microneedles for transdermal protein delivery

Microneedle mask devices have been prepared using continuous liquid interface production technique coated on polyethylene glycol base for appropriate delivery of proteins like serum albumin and ovalbumin. To avoid damage to the encapsulated protein which can eventually result in immunogenicity due to exposure to free radical polymerization procedure used to produce Continuous Liquid Interface Production-based microneedles during fabrication. The way out to this was coating microneedles after fabrication process is over. The successful delivery has been noted in studies carried out on mice for 72 hours in which sustained retention of protein was observed (Caudill et al., 2018).

3D printed Microneedles

Such types are mostly used for delivery of insulin. The cone shaped polymeric microneedles have been obtained via stereo lithography technique using a biocompatible resin. To prevent rapid degradation and maintain stability of insulin, it is coated on microneedle by using inject print technique in which insulin solution is printed on each microneedle by keeping them at
45°C and repeating the process for 92 times in which 10 dots of 2 droplets are longitudinally dispensed (Pere et al., 2018).

**Microneedles for subcutaneous tumor**

A fast dissolving microneedle patch has been fabricated using sodium hyaluronate on to which 5-aminolevulinic acid is coated to the tips of microneedle. This led to maximum drug utilization and avoidance of drug residue. Also, it helped overcome problems related with hydrophilicity and zwitterion nature of aminolevulinic acid. Hence, by using photodynamic therapy and sodium hyaluronate-based microneedles a dissolving microneedle of aminolevulinic acid proves to be best for treating subcutaneous tumor (Zhao et al., 2018).

**Patents**

The strategy of delivering drug via microneedle has resulted in several patents which are listed in Table 5. Different mechanisms for different types of microneedle were reported.

Adachi et al. (2007) developed microneedle device which helps permeate layers of skin for painless delivery of drug. Tomono (2008) utilized chitosan which is a biodegradable material for fabricating a microneedle. Lee et al. (2011) came up with an idea of developing a hermetically sealed bottom of which the microneedle array is attached at the opening of the setup of a container in which capsule kept releases drug at desired time periods.

Birchall et al. (2012) developed a system in which indicator material and therapeutic agent is coated on microneedle and the arrangements are such that drug to be delivered is delivered first and once the drug is delivered the indicator indicates the success of the drug delivered. The therapeutic agent is toward the tip and the indicator material is underneath for appropriate delivery. Herman et al. (2013) developed a system used for treating various urinary tract disorders using microneedles for drug delivery of various treatment fluids.

Levin et al. (2013) developed device that includes a syringe connected with a barrel, a plunger, and a hollow microneedle adapter. The injections, thus obtained contain microneedle adapter which are used to deliver the fluids to intradermal area. Levin, thus, devised this technique which also helps in dose sparing as it has been proven that delivering via intradermal routes produces the same efficacy as that of the doses not given in intradermal space. Vaccines are also given through this route (Levin et al., 2014).

Shaari et al. (2013) developed nasal delivery device in which certain embodiments, the nasal delivery device comprises a substrate for administration of a composition to the nasal and/or sinus mucosa, wherein the substrate is non-absorbent.

<table>
<thead>
<tr>
<th>Sr. No</th>
<th>Patent Number</th>
<th>Title</th>
<th>Abstract</th>
<th>Reference</th>
</tr>
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<tbody>
<tr>
<td>1.</td>
<td>US20070250018</td>
<td>Transdermal drug administration system with microneedle</td>
<td>Painless delivery of drug via stratum corneum to systemic circulation by using microneedle device via reservoir adsorbent and drug which gets dissolved in order to fulfill the purpose of drug delivery was patented in 2007.</td>
<td>(Adachi and Tokumoto, Higo, 2007)</td>
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<tr>
<td>2.</td>
<td>US20080200883</td>
<td>Micro-needle and micro-needle patch</td>
<td>Tomono suggested usage of biodegradable material for fabrication of microneedle and developed a successful microneedle device.</td>
<td>(Tomono, 2008)</td>
</tr>
<tr>
<td>3.</td>
<td>US20110046557</td>
<td>Microneedle drug delivery system including movable drug-containing capsule</td>
<td>Lee et al. developed movable capsules containing drug which can be delivered by microneedle drug delivery system.</td>
<td>(Lee et al., 2011)</td>
</tr>
<tr>
<td>4.</td>
<td>US20120123341</td>
<td>Monitoring system for microneedle drug delivery</td>
<td>Birchall et al. (2012) developed a monitoring system in which sequential coating of microneedle was possible with additional advantage of indications provided directly by the device regarding the completed process.</td>
<td>(Birchall et al., 2012)</td>
</tr>
<tr>
<td>5.</td>
<td>US20130331783</td>
<td>Micro-needle bladder ballon</td>
<td>Herman et al. (2013) formulated inflating balloon system using microneedles which functions to puncture the inner bladder wall for delivering particulates such as, stem cells, drugs, Biotox for treating various urinary tract disorders.</td>
<td>(Herman et al., 2013)</td>
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<td>6.</td>
<td>US20130110043</td>
<td>Microneedle intradermal drug delivery device with auto-disabled functionality</td>
<td>Levin developed an auto disable microneedle syringes for direct intradermal delivery of drugs.</td>
<td>(Levin, 2013)</td>
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<td>7.</td>
<td>US20130008547</td>
<td>Microneedle nasal delivery device</td>
<td>The invention directed by Shaari et al. lead to nasal delivery device comprising one or more microneedles, and to various methods of nasally administering a composition with a nasal delivery device comprising one or more microneedles.</td>
<td>(Shaari, 2013)</td>
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<td>9.</td>
<td>US9333330</td>
<td>Multichannel microneedle</td>
<td>Cannehan and Cachemaille (2016) developed a system with minimum two parallel independent lumens designed to open in perpendicular direction for drug delivery.</td>
<td>(Cannehan and Cachemaille, 2016)</td>
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<td>10.</td>
<td>US20160144100</td>
<td>Microneedle drug delivery system</td>
<td>Gharib et al., 2016 developed microneedle device for fluid delivery that forced drug for effective intradermal delivery or injections via needles.</td>
<td>(Gharib et al., 2016)</td>
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<td>11.</td>
<td>US20140005606</td>
<td>Embeddable microneedle patch for transdermal drug delivery and method of manufacturing the same</td>
<td>Chein and Huang invented embeddable microneedle patch which includes supporting substrate, shaft, biodegradable carrier and a drug encapsulated in the carrier.</td>
<td>(Chen and Huang, 2017)</td>
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<td>12.</td>
<td>US2017065803</td>
<td>Microneedle based cell delivery</td>
<td>Birchall et al. formulated microneedles for transplantation of cells which tends to elevate the hair growth and helps improve the skin related problems.</td>
<td>(Birchall et al., 2017)</td>
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and comprises one or more microneedles. In some embodiments, the nasal delivery device comprises a reservoir comprising one or more therapeutic agents, wherein the reservoir is in fluid communication with one or more microneedles.

Uhlrand and Peeters (2015) developed device for intraluminal delivery in which disrupted mucosal region helped in delivering of the drug through microneedles. The drug from the housing is dispersed in the disrupted mucosal region which is obtained via microneedles.

Cannan and Cachemaile (2016) lead to the development of system having independent lumen connected with elongated body, head, and sharp tip with distal opening in perpendicular direction with respect to lumen used to deliver drug via stratum corneum into systemic circulation.

Ghahrib et al. (2016) developed system using one or more needle(s) or micro-needle arrays in which description related with shift of fluid volume from a first expansion member position to a second expansion member position drives delivery of the fluid, which can be a drug or drug solution, through the needle(s) or micro-needle arrays.

Cheng and Huang (2017) developed embeddable microneedle patch in which the embeddable micro-needle patch for transdermal drug delivery is attached to the skin for a predetermined time, the biodegradable carrier is separated from the supporting shafts and embedded into the skin, and the biodegradable carrier swells and then degrade, so as to release the drugs, which are encapsulated in the biodegradable carrier, at a rate of 1%–99% loaded drug per day into the skin.

Birchall and Coulman (2017) developed microneedle device for treating skin improvement or repair comprising: a plurality of microneedles attached to or integral with a supporting base member and arranged in at least one circular pattern on same wherein said microneedles are hollow and have a bore size of between 60 and 150 μm diameter. Hence, subsequent growth can be observed in the field of microneedle drug delivery system.

CONCLUSION

Thus, this review article included the basics of transdermal drug delivery system as well as microneedle drug delivery system. Through this, one can understand that in the coming years, the importance of microneedle drug delivery is going to increase due to the advantages that it offers. It can replace the conventional methods of drug delivery most probably the transdermal approach. The biggest advantage that it offers is the painless treatment. Furthermore, the efficiency of increasing the penetration through the skin which is one of the biggest hurdles in transdermal drug delivery system has increased the scope of microneedle drug delivery system in the coming years. Also, there is no limitation of using only potent drugs for microneedle drug delivery system. Furthermore, patient compliance, possibility of self-administration, bioavailability, accuracy, precision, less chances of bacterial infections, and many other advantages leads to conclusion that the microneedle drug delivery system can definitely be used for betterment of people and eventually for good public health. Thus, microneedle drug delivery system obtained from the basis of transdermal drug delivery system can be used for better.


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