A Review on Natural Permeation Enhancer for Transdermal Drug Delivery System and Permeation Evaluation

Keywords: Transdermal drug delivery system, Permeation enhancer, Transdermal barrier, percutaneous absorption, permeabilization

ABSTRACT

The transdermal drug delivery route is evolving as a potential route due to its advantages of bypassing the hepatic first-pass metabolism, decreased side effects, and gastrointestinal effects, improve patient compliance as it is a pain-free self-administration for patients, etc. Transdermal drug delivery has been accepted as a potential non-invasive route of drug administration, with advantages of prolonged therapeutic action, decreased side effects, easy use, and better patient compliance. However, the development of transdermal products is primarily hindered by the low permeability of the skin. The major setback appearing in this route is the difficulty of the drugs to penetrate through the skin as the stratum corneum (outermost layer of the skin) forms a protective barrier for the underlying tissues from the outer environment. A transdermally delivered drug can only show its action when it can cross the transdermal barrier to reach the systemic circulation and for helping on doing that the penetration enhancer are the agents which increase the permeability of the skin which on return maintains the drug level in the blood. Permeation enhancers can be of a chemical type, natural type, and physical type. The present review describes the natural permeation enhancers can be which be employed for transdermal permeation of drugs.
INTRODUCTION:

Skin is the largest organ in the human body and its easy accessibility makes it an attractive port of entry for drug administration that dates back to the first medical records of man. The formulations that are applied to the skin can be divided into transdermal (for systemic effects) and topical (for local effects in the skin). Clinical benefits of transdermal drug delivery over conventional routes of drug administration include the following.

Figure No. 1: a) Skin layer, b) Drug properties c) Drug permeation pathways size

- sustained drug delivery over long periods (advantageous in long-term drug therapy and for patients that forget to take their drugs regularly, easy-to-check whether the drug has been taken)

- lower fluctuations in plasma levels (important for drugs with narrow therapeutic windows, may decrease the incidence of side effects or ineffective dosing)

- avoidance of metabolism and interactions in the intestine and liver (advantageous for drugs with high first-pass effect, drugs that cause nausea, interact with food and other drugs, or get inactivated in the gastrointestinal tract)

- generally good patient compliance (no needle phobia and no risk of complication from accidental needle sticks, no need to swallow pills, less frequent administration)

- ease of therapy termination (although removing the patch does not remove the drug absorbed in the upper skin layers but the concentration gradient is largely decreased)
Whilst topical formulations are widespread, transdermal drug delivery has not reached its full potential yet. The development of transdermal patches is hindered primarily by the skin’s remarkable barrier properties that evolved to keep the insides (water and electrolytes) in and outsides (chemicals, allergens, microbes) out. To enable drug administration through the skin in therapeutic doses, numerous techniques to reversibly decrease the skin barrier resistance have been developed. These methods can be broadly divided into physical and chemical approaches, and include, for example, chemical enhancers, vesicles, and particles (including high-velocity particles), microneedles, stratum corneum ablation (including lasers), iontophoresis, ultrasound, electroporation, magnetophoresis.

This review focuses on chemical permeation enhancers, also known as penetration enhancers, percutaneous absorption promoters, or accelerants. These compounds interact with the skin barrier components or alter the formulation properties to increase drug flux through the skin.

**Skin**

While scientists aiming at transdermal drug delivery may view the skin as an obstacle to their efforts, others studying the skin homeostasis in health and disease admire the multiple skin barriers, sensory properties, and self-repairing ability. This primarily protective role of the skin must be kept in mind when designing a transdermal formulation: in an attempt to deliver a drug, we should not cause irreversible damage to the skin, and we cannot expect miracles, such as the delivery of drugs that have gram daily doses. For example, with paracetamol, a simple estimation using the required plasma concentration, clearance, and human skin flux suggests that a 6 m² patch would be required.

**Dermis**

To reach cutaneous blood vessels and, subsequently, systemic circulation, a drug must get past the upper skin layer, epidermis, into the dermis. The dermis is a hydrophilic layer that does not markedly block the transport of most substances (though it can be a significant barrier for highly lipophilic compounds). Blood vessels in the dermis reach to its interface with the epidermis and remove substances that traversed the epidermal layers, maintaining the concentration gradient between the skin surface and dermis that drives the permeation. Appendages, such as hair follicles, sebaceous glands, and sweat glands, also originate in the dermis and may provide a ‘shunt’ pathway for some permeants.
Epidermis

The epidermis is composed of (from the inside to the outside) *stratum basale*, *spinosum*, *granulosum* (also collectively called the viable epidermis), and the *stratum corneum*. The cells, which are called keratinocytes, in the basal layer continuously divide and move to the surface. During this process, which is called keratinization, keratinocytes produce precursors of barrier components, such as keratin, filaggrin, and lipids that will eventually ‘seal’ the skin surface. The end product of this process, the *stratum corneum*, is the skin permeability barrier to most substances.

Stratum corneum

The *stratum corneum* consists of flattened anucleate cells, corneocytes, embedded in a lipid matrix. Corneocytes are filled with keratin bundles and other proteins such as filaggrin, and their cell membrane is replaced by a highly cross-linked protein layer, termed corneocyte (cornified) envelope. The corneocyte envelope is further decorated with a covalently attached monolayer of ultralong lipids, called corneocyte lipid envelope. This lipid monolayer likely acts as a scaffold for the orientation of the extracellular lipid matrix and prevents a permeable boundary between the hydrophilic cells and hydrophobic extracellular domains.

![Figure No. 2: Mechanism of action of permeation enhancers](image-url)
Stratum corneum lipids

The extracellular lipid matrix in the stratum corneum is unique in its composition and organization. The major stratum corneum lipid classes comprise ceramides, free fatty acids, and cholesterol in an approximately 1:1:1 molar ratio; phospholipids are virtually absent. Notably, the skin ceramides are extremely heterogeneous – over 1000 distinct structures have been identified so far. This heterogeneity probably provides a ‘buffering’ capacity of these barrier lipids so that they can adapt to environmental changes and/or incorporate foreign substances (to a certain extent) without significantly compromising the barrier integrity. Ceramides are simple sphingolipids (sphingoid bases with a fatty acid attached to the amino group); thus, their common features include a small polar head that does not extensively hydrate and two hydrophobic chains (mostly saturated) of unequal lengths – the sphingoid base chain usually has 18 carbons whereas the acyl chain mostly reaches 24 carbons (very-long-chain) or over 30 carbons (ultralong chain). These lipids assemble into tightly packed multilayered lamellae of 6 nm and, in particular, 13 nm repeat distances.

Transdermal drug delivery offers a very advantageous route for drug delivery compared with the other routes of drug administration having advantages such as bypassing the hepatic first-pass metabolism, and longer duration of action. However, the barrier function of the skin outermost layer, the stratum corneum (SC) is one of the main limitations to it, and for this reason, skin penetration enhancers are gaining the greatest interest in pharmaceutical research. Penetration enhancers help in the permeation of the desired drug (penetrant) through the skin by lowering the impermeability of the skin. Some properties which are desired in permeation enhancers are the must be pharmacologically inert, nonirritating, nontoxic, nonallergic, compatible with drugs and excipients, odorless, tasteless, colorless, and inexpensive, and also have good solvent properties. For a transdermal matrix patch to be successful, it relies on the capability of the penetrant (drug) to penetrate the skin in quantities which is sufficient to maintain therapeutic levels. During formulating transdermal drug delivery system (TDDS), chemical enhancers are used widely as it helps in reducing the permeation barrier properties of the skin. Several literature reviews mentioned that naturally occurring oils (essential oil and vegetable oil) can be employed as permeation enhancers due to their advantageous properties such as compatibility with drug and excipients, nontoxic, less allergic, and easy availability. This review article describes the barrier properties of the skin and various natural penetration enhancers that have been employed to enhance the transdermal permeation of drugs and the different parameters for permeation evaluation.
SKIN AS A BARRIER TO DRUG PERMEATION

The outermost few microns of the skin, the SC, aids to the barrier function of the skin. This layer of the skin is the most impermeable, forming a laminate of compressed keratin-filled corneocytes attached in a lipophilic matrix. The lipids of this matrix are distinguishing in many respects: • From the skin surface to the base of the SC, they provide the only continuous phase • Among the biomembranes, the absence of phospholipid is particular and is a composition (ceramides, free fatty acids, and cholesterol) that is unique • The SC lipids exist as multilamellar sheets even though it is having a deficit of polar bilayer forming lipids • The essentially saturated, long-chain hydrocarbon tail aids a highly ordered, interlock configuration. However, the resistivity of the membrane cannot be entirely explained by the unusual lipid matrix, and the architecture of the SC altogether has been proposed to play a role in the barrier property of the membrane. The corneocyte resembling a brick and mortar assembly is suggested to impart the membrane-impermeable to water concerning other biomembranes. And by the visualization studies localizing several permeants, in the intercellular channels by kinetic analysis of the in vivo skin penetration rates of model compounds and by evidence from thermotropic biophysical studies of lipid domains the transport role of this pathway gets furthermore support.

Transdermal permeation pathways

Transdermal permeation can take place by diffusion through A. Through the SC: Using the transcellular route, drugs can pass through the corneocytes highly hydrated keratins are present in corneocyte, which provides an aqueous environment for which the hydrophilic drugs can pass. Hence, the transcellular pathway is a predominant pathway for hydrophilic drugs B. Intercellular permeation: In the intercellular pathway, the drug diffusing takes place utilizing the continuous lipid matrix C. Transappendaged permeation: Only 0.1% of the skin surface area is covered by the hair follicle and sweat glands which limit the area available for the applied drug formulation to come in contact with it. For many drugs, an aqueous pathway is considered desirable but as the sweat is traveling against the diffusion pathway of the permeant, permeation may be limited. Lipid-rich sebum fills the sebaceous gland, which may present a barrier for hydrophilic drugs.
PHYSICOCHEMICAL ASPECTS OF SKIN PENETRATION

The diffusion of the drug through the skin is a passive kinetic process that occurs down the concentration gradient (from a high concentration region to a low concentration region). The steady-state equation can be described by Fick’s first law of diffusion. The equation describes the rate of transfer (flux, J) of a diffusing substance through the unit area of the membrane and diffusion coefficient, D to the concentration gradient across the membrane (dc/dx). J=−AD (dc/dx) (1) The negative sign in equation (1) is because the diffusion process occurs in the opposite direction to increased concentration. Fick’s second law of diffusion, equation (2) can be derived from equation (1) to describe membrane transport under nonsteady state condition: dc/dt=D d2c/dx2 (2) By maintaining the sink conditions in the receptor compartment and maximum fixed concentration in the donor compartment, the equation (2) can be written as J=AD (Cm/ɦ) (3) Where Cm is the concentration in the donor-membrane interphase and ɦ is the effective diffusional path length. The Cm in the equation (3) can be replaced by vehicle membrane partition coefficient (K) as the ratio between the concentration of permeant in the membrane at the donor-membrane interface and the vehicle in which applied (Cv). Modified Fick’s first law of diffusion describes the steady-state flux across the membrane equation. J SS=ADKCv/ɦ (4) We can conclude that increased drug flux can be achieved by a change in D, K, and C. The compounds which are skin penetration enhancers should potentially change the solubility or partition behavior of the drug into SC or its diffusion properties or both sometimes change in the thermodynamic activity of drugs in the formulation manipulate the flux.

PERMEATION ENHANCERS

Permeation enhancers are compounds that promote skin permeability. They are an important factor in a TDDS which is used to improve the flux (J). Flux can be defined as the amount of material flowing through the unit cross-section area at the time (t). Ideal properties of penetration enhancers: • They must be pharmacologically inert, nonallergic, nonirritating, and nontoxic • It must have compatibility with excipients and drugs • It must not have any pharmacological activity in the body • Cosmetically it must be acceptable • It must be odorless, tasteless, and colorless • It should allow therapeutic agents into the body but should prevent the loss of endogenous material from the body, i.e., they should work unidirectionally • It must have chemical and physical stability • It must have a reproducible and predictable duration of action • It must have a good solvent property.
NATURAL PERMEATION ENHANCERS (NPEs)

NPEs are a comparatively new class of penetration enhancers in the pharmaceutical industry. Due to its advantages such as low cost, better safety profile more research needs to be focused in this field to develop stable transdermal formulations containing natural permeation enhancers (NPEs) which can be scaled up for commercial transdermal drug product.

Papain

Papain is isolated from Carica papaya. It is an endocytic plant cysteine protease enzyme.

Papain, which is a proteolytic enzyme, was studied in vitro and in vivo permeation of low-molecular-weight heparin (LMWH). The combined administration of LMWH and papain was found to be a new approach in improvement in the absorption of orally administered heparin and hence its bioavailability.

Piperine

Piperine is obtained from the mature fruits of Piper nigrum and Piper longum. Piperine was investigated for in vitro permeation of aceclofenac across human cadaver skin, and Fourier transforms infrared technology was used to check the possible mechanism from which the results obtained showed that piperine enhances transdermal permeation of aceclofenac by a biphasic mechanism involving partial extraction of SC lipid and interaction with SC keratin.

Capsaicin

Capsaicin is a major alkaloid among capsaicinoids which is produced only in capsicum fruits of the genus capsicum and belonging to the Solanaceae family. The permeation enhancing properties of capsaicin were studied for naproxen taking azone as the standard enhancer and capsaicin were compared to it. Onto the skin, different amount of chosen enhancer was applied before the experiment. A formulation containing 3% capsaicin and commercially available naproxen gel formulation was also studied, and the results were compared. It was found that penetration increased when the skin was treated with azone and capsaicin and also some alteration by capsaicin was seen in the SC layer. Hence, it was observed that capsaicin increases the penetration of naproxen through SC, which concludes that capsaicin is the quiet capable enhancer of skin like well-known enhancer azone.
**Myristica fragrans**

*M. fragrans* was evaluated as a penetration enhancer in a transdermal gel formulation containing diclofenac sodium as the target drug. Methanolic extracts, chloroform extracts, and n-hexane extracts of *M. fragrans* were used as a penetration enhancer in comparison to a synthetic enhancer Triton X. It was found that in both in vivo and in vitro studies, methanol and chloroform extract showed better percentage cumulative release (%), and hence, better penetration was shown as compared to the synthetic enhancer.

**Novel permeation enhancers**

**Ceramide analogs**

The lipids in the SC consist of ceramides, cholesterol, and fatty acids, as well as a small quantity of cholesterol sulfate. It has been known that ceramides are the main components responsible for the barrier effect of the SC. Structurally, they are amphiphilic molecules, containing two long hydrocarbon tails and an acylamide polar head. Vávrová et al. believed that certain structure similarity might exist between enhancers and ceramides so that the enhancer molecules could insert themselves between the hydrophobic tails of the ceramide bilayers and weaken the continuity of the lipid barrier. In their study, a series of ceramide analogs with two kinds of polar heads based on L-serine and glycine were respectively synthesized, and the relationship between the properties of the polar heads and enhancement activity was discussed. The result indicated that although these compounds were inactive in isopropyl myristate (IPM), significant enhancement effects were observed when they were used in the water (W) suspensions. Especially, compound 1 with the simplest glycine group as the polar head structure was the most potent one, and its promoting activity was significantly higher than compound 2, an L-serine based derivative which had a hydroxyl group in the polar head. Furthermore, these two compounds were both more active than their respective analogs with dipeptide as the polar heads, which were shown by compound 4 and 5. These results led to a hypothesis that the increasing size and H-bonding ability would decrease the activity of the ceramide analogs, and it might account for the little enhancing activity of compound 6, which possessed a large polar head with three hydroxyl groups. Also, the study suggested that the introduction of a double bond into the hydrocarbon chain did not contribute much to the activity of the present ceramide analog, which was demonstrated by the compound.
Azone analogs

Azone (1-dodecylazacycloheptan-2-one or laurocapram, compound, is the first molecule specially designed as a skin permeation enhancer. It has been demonstrated to enhance the transdermal absorption of a wide variety of drugs. Azone probably exerts its permeation enhancing effects mainly via the interaction with the lipid domains of the SC. The ‘soup spoon’ structure would make it more likely to partition into the bilayer lipids, where Azone might exist dispersed or in separated domains to disrupt the organized lipid packing. Although the FDA has not approved its use in pharmaceutical products for its slight side effects, Azone is recorded in the Chinese Pharmacopoeia and widely used in China. To improve its properties, hundreds of Azone like compounds have been synthesized, which provide a huge amount of information about the molecular design for Azone analogs.

Essential Oil

Essential oil is natural products that are extracted from aromatic plants having a concoction of several aromatic smelling volatile compounds, primarily consisting compounds such as terpenes, terpenoids, and phenylpropanoids [21]. They can be accepted as a natural alternative to synthetic skin penetration enhancer due to their promising penetration enhancing activity [22]. As a penetration enhancer, essential oils help in the delivery of drug compounds into the skin by interacting with the intercellular lipids by different physical processes such as increased disorder, phase separation, and fluidization. As they are penetrated easily by the skin, they are easily excreted also by the body with urine and feces. Hence, due to their better safety profile in comparison with other penetration enhancers, their use is increasing.

Essential oil as a skin permeation enhancer

Penetration enhancer interacts with the tissue components to lessen the barrier properties by partitioning into the SC but do not cause any damage to the underlying skin cell. Both D-limonene and 1, 8-cineole have been shown to modify permeant diffusivity by disrupting SC lipid.

Eucalyptus oil

The oil of eucalyptus can be obtained from several species of the Myrtaceae family, such as Eucalyptus citriodora, Eucalyptus dives, Eucalyptus globulus, Eucalyptus polybractea, and
**Eucalyptus radiata.** By steam distillation of eucalyptus leaves, its oil is extracted. The eucalyptus oil was subjected to permeation studies on full-thickness human skin and it was found that the oil enhanced the permeation of chlorhexidine (2% [w/v]) into the dermis and the lower layer of the epidermis when it was combined with 70% (w/v) isopropyl alcohol and 10% (v/v) eucalyptus oil in comparison to the solution of chlorhexidine/isopropyl alcohol alone.

**Niaouli oil**

The extraction of niaouli oil can be made by steam distillation of twigs and leaves of *Melaleuca quinquenervia*, of the family Myrtaceae. The major constituents of niaouli oil was 55-70% 1,8-cineole (oxide) and limonene (monoterpene), 7-15% a-pinene (monoterpene), 2-6% s-pinene (monoterpene), and 2-6% viridiflorol (sesquiterpene).

In vitro studies were performed to determine the permeation enhancing the effect of niaouli oil at 10% (w/w) concentration in propylene glycol on estradiol model drug using a hairless mouse skin. It was found that niaouli oil proved to be more effective in transdermal permeation of estradiol than cajput, myrtle, orange, and cardamom essential oil.

**Fennel oil**

The extraction of fennel oil can be made from the seeds of *Foeniculum vulgare*, of the family Umbelliferae. On permeation studies, the percutaneous penetration of trazodone hydrochloride was enhanced by fennel oil followed by eucalyptus oil, citronella oil, and mentha oil. The variable physicochemical properties and molecular weights of phytochemicals present in the different essential oils may be the factors for the differences in the permeation enhancement activity between the oils.

**Black cumin oil**

The extraction of black cumin oil is made by steam distillation of the seeds of *Cuminum cyminum* by steam distillation. Black cumin oil showed a relatively greater permeating effect for the drug carvedilol when it was compared to clove oil, eucalyptus oil, tulsi oil, oleic acid, Tween 80 and the enhancement factor was found to be 6.40. Furthermore, Fourier transform infrared spectroscopy studies confirmed the alteration caused by black cumin oil on the permeability of the skin by extracting lipids and by hydrogen bonding which affects other hydrogen bonds between the ceramides.
Almond oil

Oil of almond and oleic acid were found as promising carriers/vehicles for enhanced permeability and solubility of aceclofenac. Hence, these oils can be used to develop drug delivery systems for improved bioavailability of aceclofenac.

In the study, topically applied ketoprofen gels and patches were formulated and evaluated, and almond oil was checked as a penetration enhancer for ketoprofen gels and patches through artificial membrane/ rabbit skin. It was found that almond oil in different concentrations as a penetration enhancer enhances the penetration of drugs from transdermal gels and patches across synthetic membrane/rabbit skin but notably when used in 3% concentration.

Basil oil

Basil oil was studied for its potential as a permeation enhancer for labetalol hydrochloride concerning camphor, thymol, geraniol, and clove oil. It was suggested that basil oil is having good penetration enhancing property for improved transdermal drug delivery of labetalol. Basil oil was used for the enhancement in the bioavailability of flurbiprofen applied transdermally and it was concluded that the bioavailability of the flurbiprofen applied transdermally using basil oil increased by 2.97, 3.80, and 5.56 times in comparison to the flurbiprofen administered orally in albino rats.

Alpinia oxyphylla oil

*Alpinia oxyphylla* oil was extracted from A. oxyphylla and was divided into a higher polarity fraction and a lower polarity fraction. In vitro studies were performed using Franz diffusion cell across the dorsal skin of Wistar rats, the results indicated that the high polarity fraction of A. oxyphylla oil was having more efficient permeation enhancing the effect of indomethacin at concentration 3% and 5% then the lower polarity fraction.

Turpentine oil

Turpentine oil on the skin permeation rate of flurbiprofen showed an additive effect when it was added to an optimized cosolvent mixture of propylene glycol and isopropyl alcohol (30-705 [v/v]), and at the concentration of 5% (v/v) of turpentine oil, the maximum transdermal penetration rate was obtained. The effectiveness of turpentine oil was investigated for permeation enhancing activity for diclofenac dimethylamine matrix patches across the
artificial skin in the Franz diffusion cell. It was found that the oil showed increasing permeation with increasing concentration of turpentine.

**Rosemary oil**

Rosemary oil is extracted from Rosmarinus officinalis. Rosemary oil when investigated for skin permeation enhancing activity for diclofenac sodium topical gel showed enhanced skin absorption at 0.5% and 1% concentration, respectively.

**Cardamom oil**

Cardamom (Elettaria cardamomum) is a common spice of India belonging to the Zingiberaceae family. The oil extracted from cardamom has several volatile compounds such as monoterpenes including 1,8-cineole and cis-ocimene and sesquiterpene including guanine and nerodilol. Cardamom oil on in vitro permeation studies through the rabbit abdominal skin showed an increase in penetration of the drugs indomethacin, diclofenac, and piroxicam.

**TERPENES**

In transdermal drug delivery studies, terpenes are one of the major choices. This class includes a heterogeneous range of members. The physicochemical properties of a specific terpene play a role in showing the effect on the skin, in particular, its lipophilicity. However, smaller terpenes with nonpolar groups are said to be better skin permeation enhancers. Terpenes are also reported to increase drug partitioning and diffusibility into the skin by disturbing the lipid bilayers of the skin. They are relatively safe as skin penetration enhancers for both hydrophilic and lipophilic drugs. Farnesol is present in many essential oils, such as citronella, neroli, cyclamen, lemongrass, tuberose, balsam, and tolu. It is sesquiterpene alcohol. Farnesol (0.25%) was reported to increase the permeation of diclofenac sodium concerning other terpenes in the following order: Farnesol > carvone > nerolidol > menthone > limonene oxide.

**Menthol**

Menthol, which is one of the potent penetration enhancers, is obtained from the flowering tops of *Mentha piperita*. Menthol and limonene together can be used as a prototype of terpenes that can be used as permeation enhancers. Eucalyptol is a cyclic ether and a monoterpenoid known by several synonyms such as 1, 8-cineole, cajeputol, eucalyptol, and cineole. Eucalyptol is used in cosmetics, fragrances, and flavoring industries because of its
spicy aroma and taste. 1, 8-cineole has also been used for the percutaneous absorption of several lipophilic drugs through the hairless mouse skin.

Eugenol was evaluated for permeation enhancing effect for the drug lornoxicam which is a nonsteroidal anti-inflammatory drug of oxicam class. Lornoxicam transdermal patches were formulated, then subjected to in vitro studies in a Franz diffusion cell using rat skin. In vitro studies showed that eugenol does increase the permeation of lornoxicam across rat skin. Borneol The transdermal permeation enhancing activity of borneol was investigated on 5 model drug, namely, 5-fluorouracil, antipyrine, aspirin, salicylic acid, and ibuprofen and it was found that borneol effectively promoted the transdermal permeation of the model drugs.

PERMEATION STUDIES

The estimation of percutaneous absorption of a molecule is a very important step in the evaluation of a molecule of TDDS. Moreover, in both pharmaceutical and cosmetic research, to support the development of TDDS skin permeation rates, evaluation has become a very important step. The various ways by which the transdermal permeation of a drug molecule can be studied are. In vitro release studies Franz diffusion cell is used for carrying out in vitro release studies wherein the donor compartment the circular patch is mounted, the receptor compartment of which is filled with phosphate buffer saline (pH - 7.4), and the whole assembly is maintained at 32°C by the warm water circulation through the water jacket and is stirred at 40-50 rpm. The patch was kept in contact with the receptor liquid. About 0.5 ml of the sample was withdrawn at regular intervals for 8 hrs and was replaced with the same amount of the medium. The samples were subjected to filtration, diluted, and analyzed spectrophotometrically. In-vivo release studies Animal models Hairless rat, hairless rhesus monkey, guinea pig, and rabbit are the most commonly used species used for evaluating the transdermal drug. One of the most reliable models for in vivo evaluation of transdermal delivery in man is the rhesus monkey. Animals on small scale are preferred for in vivo studies because of economic factors and availability. Human volunteers Clinical phase is one of the important stages in the development of a transdermal device, and in this stage, the formulation is tested in human volunteers for the collection of pharmacokinetic and pharmacodynamics data which are required to evaluate the generation of toxic effect during application of the formulation. C14 radioisotope is used for labeling drugs to determine percutaneous absorption in humans and measuring the radioactivity in excreta, but for knowing, the amount residing in a body and amount excreted by other route attention is
required. In vivo dermal absorption study For in vivo dermal absorption test, the dorsal surface (approximately 4 cm × 3 cm area) of the rabbit skin is first clipped free for the application of the test substance and this is done before 24 hrs of starting the test. The requisite amount of the test substance was applied to the area, and 1 ml of the blood sample was withdrawn from the central ear artery at predetermined times over 24 hrs at 2, 6, 12, 24 hrs. Plasma was obtained by centrifuging the blood and was stored at −20°C before analysis. The analysis was done using headspace gas chromatography-mass spectrometry.

CONCLUSION:

The field of transdermal drug delivery due to its advantages has been rapidly developing and has stimulated various researches to incorporate more and more drugs through the transdermal route. Skin serves a limitation for permeation of drugs, and hence, permeation enhancers are used to increase the permeability of the poorly absorbed drugs and hence maintain its bioavailability. The review article summarizes the various NPEs which can be used to accelerate the permeation of drug across the skin for the development of the transdermal delivery system and the parameters by which the permeation studies can be done. In this article, various NPEs has been discussed which can be used to accelerate the permeation of drug across the skin for the development of TDDS.

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