Fast Dissolving Films: An Innovative Drug Delivery System

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Abstract

In the recent years, many of the pharmaceutical groups are focusing their research on rapid dissolving technology. Amongst the plethora of avenues explored for rapid drug releasing product, Fast Dissolving Films technology is gaining much attention. Fast Dissolving Films evolved over the past few years from the confection and oral care markets in the form of breath strips and became a novel and widely accepted form by consumers for delivering vitamins and personal care products. These are solid dosage forms, which disintegrate or dissolve within 1 min when placed in the mouth without drinking water or chewing. This technology has been used for local action, rapid release products. The oral films are formulated using polymers, plasticizer, flavors, colors and sweeteners. This review describes about the formulation methodology, evaluation parameters and the future aspects of fast dissolving films.

Keywords: Fast Dissolving Films, Solvent casting technique, Rapid disintegrating, Patented technologies

Introduction

Oral route of administration is the most convenient and preferred route of administration among the various other delivery system. More than 70% of drugs are available in the market in the form of oral drug delivery system due to pain avoidance and versatility (to accommodate various types of drug candidates) [1]. Dysphagia is commonly found among all age groups. Due to this problem, approximately 50% of population, mainly pediatric and geriatric patients, tend to avoid taking oral solid dosage preparations due to fear of choking. To overcome various problems related to swallowing, Fast dissolving Tablets (FDTs) were designed in early 19th century, which slowly led to their further advancement and thus Fast Dissolving Films (FDFs) were developed. Fast dissolving dosage form has become increasingly important because of their unique properties. They quickly disintegrate and dissolve, and can be administered without water, making them particularly suitable for pediatrics and geriatric patients. Fast dissolving films (FDFs), have gained popularity not only in breath strips but also in personal care, food and drug delivery markets [2,3]. Pharmaceutical companies and consumers alike have embraced FDFs as a practical and accepted alternative to traditional OTC medicines, such as liquids, tablets and capsules, because of the various benefits of the films. FDFs offer fast, accurate dosing in a safe, efficacious format that is convenient and portable, without requiring the use of water or a spoon. A variety of polymers are available for preparation of RDFs. The polymers can be used alone or in combination to obtain the desired film properties. The films obtained should be tough enough so that they are not damaged while handling or during transportation. On the other hand, Mouth dissolving films should have the property to dissolve within seconds when placed in mouth and deliver the drug to the oral cavity instantaneously. The excipients employed in FDFs are preferably hydrophilic in nature whereas drug may be either hydrophilic or hydrophobic. As the film forming polymer (which forms the platform for the RDFs) and plasticizer are the most essential and major component of the FDFs, at least 40-50 % w/w of polymer and upto 20% (total weight of...
polymer) of plasticizer should generally be present based on the total weight of dry RDFs. Technology Catalysts forecasts the market for drug products in oral thin film formulations was valued of $500 million in 2007 and could reach $2billion in 2012. Based on upward global growth trends of the past decade, the fast dissolving dosage market could produce revenues of $13billion by 2015[4]. The FDFs technology continues to be viewed as an alternative for FDT products that would afford a superior barrier to generic entry and product differentiation to over-the-counter brands. From the marketing perspective, a patented ODF technology would be beneficial. The grant of marketing exclusivity to the new dosage form would help to gain more revenue. The various synonyms used for FDFs include mouth dissolving films (MDFs), orally disintegrating films (ODFs), melt in-mouth films, oro-dispersible, quick dissolving and rapid disintegrating films.[4,5]

Advantages of Fast Dissolving Films [6]
• No risk of choking and obstruction.
• No need of water has led to better acceptability amongst the dysphagic patients
• Improved oral bioavailability of drugs
• Taste masking
• Enhanced stability
• Improved patient compliance
• Oral films are flexible and they are not as fragile as most of the ODTs
• Reduction in first pass metabolism may lead to reduction in the dose
• The oral or buccal mucosa is highly vascularized, hence drugs can be absorbed directly and can enter the systemic circulation without undergoing first-pass hepatic metabolism

Ideal Properties of Fast Dissolving Films
• It should have an acceptable taste.
• It should give a pleasing mouth feel.
• It should be less friable and have good mechanical strength to withstand the post manufacturing handling.
• It should be stable in environmental conditions.
• Subsequent to oral administration, it should leave least or no residue in mouth.
• It should quickly dissolve to release drug instantaneously in mouth.
• It should be compatible with the other ingredients.

• The drug should have pleasant taste.
• The drug to be incorporated should have low dose upto 40 mg.
• The drugs with smaller and moderate molecular weight are preferable.
• Good solubility in water as well as in saliva and also good stability.

Classification of Oral Films[6]
There are three different subtypes of oral films:

i. Flash release
ii. Mucoadhesive melt-away wafer
iii. Mucoadhesive sustained-release wafers

Types of oral films and their properties are described in Table 1

Classification of Fast Dissolving Technology
For ease of description, fast-dissolve technologies can be divided into three broad groups:

i. Lyophilized systems: The technology around these systems involves taking a suspension or solution of drug with other structural excipients, through the use of a mould or blister pack, forming tablet-shaped units. The units or tablets are then frozen and lyophilized in the pack or mould. The resulting units have a very high porosity, which allows rapid water or saliva penetration and very rapid disintegration.

ii. Compressed tablet-based systems: This system is produced using standard tablet technology by direct compression of excipients. Depending on the method of manufacture, the tablet technologies have different levels of hardness and friability. The speed of disintegration for fast-dissolve tablets compared with a standard tablet is achieved by formulating using either water soluble excipients, superdisintegrant or effervescent components, to allow rapid penetration of water into the core of the tablet.

iii. Thin film strips: Oral films, also called oral wafers, evolved over the past few years from the confection and oral care markets in the form of breath strips and became a novel and widely accepted form by consumers for delivering vitamins and personal care products. Today, FDFs are a proven and accepted technology for the systemic delivery of APIs for over-the-counter (OTC) medications and are in the early- to mid development stages for prescription drugs. This has been attributed to the success of the breath freshener products by consumers such as Listerine Pocket Paks in the US consumer
market. Such systems use a variety of hydrophilic polymers to produce a 50-200 mm film. The film is manufactured as a large sheet and then cut into individual dosage units for packaging in a range of pharmaceutically acceptable formats.


1) Drug Category
2) Film Forming Polymers
3) Plasticizers
4) Sweetening Agents
5) Saliva Stimulating Agents
6) Cooling Agent
7) Flavoring Agent
8) Coloring Agent
9) Surfactants
10) Stabilizing and thickening agents

Formulation of FDFs involves the intricate application of aesthetic and performance characteristics such as taste masking, fast dissolution, physical appearance, mouth feel etc. From the regulatory perspectives, all excipients used in the formulation of OS should be Generally Regarded as Safe (i.e. GRAS-listed) and should be approved for use in oral pharmaceutical dosage forms. A typical composition includes various ingredients which are described in the Table 2.

i. Drug Category: This technology has the potential for delivery of variety of APIs. However since the size of the dosage form has limitation, high dose drugs are difficult to be incorporated in films. Several classes of drugs can be formulated as fast dissolving films including antiulcer, antiasthamatics, antitussives, expectorants, antihistaminics, NSAID’S etc.

ii. Film Forming Polymers: Water-soluble polymers are used as film formers as they provide rapid disintegration, good mouthfeel, and mechanical strength to the films. The robustness of the strip depends on the type of polymer and its amount in the formulations. Water-soluble polymers film, adheres to the buccal mucosa and rapidly delivers medication into the systemic circulation. A variety of polymers are available for preparation of films of which pullulan, gelatin and hypropomellose are most commonly used. At least 45% w/w of polymer should generally be present based on the total weight of dry film [18]. Examples of water-soluble polymers include: Pullulan, Gelatin, guar gum, Xanthum gum, Hydroxyl propyl methyl cellulose, Modified starches, Hydroxyl ethyl cellulose etc.

iii. Plasticizers: Plasticizer is a vital ingredient of the oral films. The selection of plasticizer depends upon its compatibility with the polymer and also the type of solvent employed in the casting of film. It helps to improve the flexibility of the film and reduces the brittleness of the film. Plasticizer significantly improves the strip properties by reducing the glass transition temperature of the polymer. Typically the plasticizers are used in the concentration of 1 - 20% w/w of dry polymer weight. Examples include: Glycerol, Propylene glycol, Low molecular weight polyethylene glycols, Citrate derivatives like triacetin, acetyl citrate, Phthalate derivatives like dimethyl, diethyl, dibutyl derivatives, Castor oil etc.

iv. Sweetening agents: Sweeteners have become the important part of the food products as well as pharmaceutical products intended to be disintegrated or dissolved in the oral cavity. The sweet taste in formulation is more important in case of pediatric population. Natural sweeteners as well as artificial sweeteners are used to improve the palatability of the mouth dissolving formulations. Suitable sweeteners include:

(a) Water soluble natural sweetener: xylose, ribose, glucose, sucrose, maltose, stevioside etc.
(b) Water soluble artificial sweetener: sodium or calcium saccharin salts, cyclamate salts, acesulfame-k etc.
(c) Dipeptide based sweetener: aspartame

v. Saliva stimulating agent: The purpose of using saliva stimulating agents is to increase the rate of production of saliva that would aid in the faster dissolution of the film formulations. Generally acids which are used in the preparation of food can be utilized as salivary stimulants. Citric acid, malic acid, lactic acid, ascorbic acid and tartaric acid are the few examples of salivary stimulants, citric acid being the most preferred amongst them.

vi. Cooling agents: Cooling agents like monomethyl succinate can be added to improve the flavor strength and to enhance the mouth-feel effect of the product. Other cooling agents like WS3, WS23 and Utracoll II can also be used in conjunction with flavors [31].

vii. Flavoring agents: Perception for the flavour changes from individual to individual depending on the ethnicity and liking. It was observed that age plays a significant role in the taste fondness. Flavoring agents can be selected from synthetic flavor oils, oleo resins, extract derived from various parts of the plants like leaves, fruits and flowers. Peppermint oil, cinnamon oil, oil of nutmeg are examples of flavor oils while vanilla, cocoa, coffee, chocolate and citrus are fruity
flavors. Apple, raspberry, cherry, pineapple are few examples of fruit essence type. The amount of flavor needed to mask the taste depends on the flavor type and its strength.

**viii. Coloring agents:** Pigments such as titanium dioxide or FD&C approved coloring agents are incorporated (not exceeding concentration levels of 1% w/w) in OS when some of the formulation ingredients or drugs are present in insoluble or suspension form.

**ix. Surfactants:** Surfactants are used as solubilizing or wetting or dispersing agents so that the film gets dissolved within seconds and release active agent immediately. Surfactants also improve the solubility of poorly soluble drugs in fast dissolving buccal films. Some of the commonly used are poloxamer 407, sodium lauryl sulfate, benzalkonium chloride, benzthonium chloride, tweens and spans etc.

**x. Stabilizing and thickening agents:** The stabilizing and thickening agents are employed to improve the viscosity and consistency of dispersion or solution of the strip preparation solution or suspension before casting. Natural gums like xanthan gum, locust bean gum, carragenan and cellulose derivatives can be used in the concentration up to 5% w/w as thickening agents and stabilizing agents.

**Methods of Manufacturing Fast Dissolving Films**

Following are the methods of manufacturing for fast dissolving films. One or combination of the following process can be used to manufacture the fast dissolving films –

i. Solvent casting method
ii. Semisolid casting method
iii. Hot melt extrusion
iv. Solid dispersion extrusion
v. Rolling method

Generally the solvent casting method is employed for manufacture of strips.

1) Solvent Casting Technique

Fast dissolving films are preferably formulated using the solvent casting method, whereby the water soluble ingredients are dissolved to form a clear viscous solution and the drug along with other excipients is dissolved in suitable solvent then both the solutions are mixed and finally casted in to the Petri plate and dried, which is then cut into pieces of the desired size. The properties of the API play a critical role in the selection of a suitable solvent. Water-soluble hydrocolloids used to prepare RDFs include: hydroxyl propyl methyl cellulose (HPMC), hydroxyl propyl cellulose (HPC), pullulan, sodium alginate, pectin, carboxy methyl cellulose (CMC), polyvinyl alcohol (PVA). Solvents used for the preparation of solution or suspension should ideally be selected from ICH Class 3 solvent list [2]. Specific types of equipment such as rollers are required for pouring the solution on an inert base. The clearance between the roller and the substrate determines the required thickness of the film. The final step, drying the film, removes the solvent and helps to obtain the finished product. Usually, glass, plastic, or teflon plates are used as an inert base for film casting. When the manufacturing technology is transferred from laboratory scale to production scale, several problems can be encountered. These problems can include the casting of the film, obtaining uniform thickness of the film, and proper drying of the sample. The selection of the proper type of dryer is needed in the final step of drying.

Once the films are dried, cutting, stripping, and packaging is done. Suitable size and shapes of films can be cut. The commonly available sizes of films are 3 x 2 cm² and 2 x 2 cm². Flowchart showing the solvent casting method is described in figure.1.

2) Semisolid casting

In semisolid casting method, firstly a solution of water soluble film forming polymer is prepared. The resulting solution is added to a solution of acid insoluble polymer (e.g. cellulose acetate phthalate, cellulose acetate butyrate), which can be prepared in ammonium or sodium hydroxide. Then appropriate amount of plasticizer is added so that a gel mass is obtained. Finally the gel mass is casted in to the films or ribbons using heat controlled drums. The thickness of the film is about 0.015-0.05 inches. The ratio of the acid insoluble polymer to film forming polymer should be 1:4.

3) Hot melt extrusion

Hot melt extrusion is commonly used to prepare granules, sustained-release tablets, and transdermal and transmucosal drug-delivery systems. In hot melt extrusion method firstly the drug is mixed with carriers in solid form. Then the extruder having heaters melts the mixture. Finally the melt is shaped in to films by the dies. Usually, when designing RDFs, polymers with low molecular weight or viscosity, such as HPMEC E5 or pullulan PI.20, are preferred. A combination of various grades of polymers may also be used to achieve desired physical properties. Mixing polymers of high and low viscosity produces a film with good mechanical strength and high drug solubility in the film. The manufacturing process for the wafers in the pharmaceutical industry is divided into different steps.
Generally, the mass is prepared first under the control of temperature and steering speed. Afterwards, the wafers are coated and dried in a drying tunnel, once again the temperature, air circulation and line speed are controlled. Then follows a slitting and in the last step the wafers are punched, pouched and sealed. Other ways of manufacturing oral wafers are spraying process or extrusion, in particular hot-melt extrusion. There are certain benefits of hot melt extrusion:

- Fewer operation units
- Better content uniformity
- An anhydrous process

Disadvantages:
- Thermal process so drug/polymer stability problem
- Flow properties of the polymer are essential to processing
- Limited number of available polymers

Comparison of solvent-casting and HME for the manufacturing of ODFs is described in the Table 3.

5) Solid dispersion extrusion
The term solid dispersion refers to the dispersion of one or more APIs in an inert carrier in a solid state in the presence of amorphous hydrophilic polymers using methods such as HME. In this method, immiscible components are extruded with drug and then solid dispersions are prepared. Finally the solid dispersions are shaped in to films by means of dies.

6) Rolling Method
In rolling method a solution or suspension containing drug is rolled on a carrier. The solvent is mainly water or a mixture of water and alcohol. The film is dried on the rollers and cutted into desired shapes and sizes. Three roll coating unit Diagram is shown in fig 2.

Objective of Formulation of Fdf's
The aim of the present research work is development and characterization of mouth dissolving oral films of a suitable drug candidate so as to achieve following objectives:

i. To improve patient compliance.
ii. To provide a rapid onset of action.
iii. To reduce the extent of hepatic first pass metabolism.
iv. To reduce the dose administered and thus the side effects associated with it.
v. To enhance the oral bioavailability of molecules.

Patented Technologies [13]

1) XGel: XGel is at the heart of Meldex international’s intellectual properties used in all its film system and its ingestible delivery technologies. XGel film Technology developed by BioProgress is bringing a revolution in the product offerings and manufacturing methods now available to the pharmaceutical industry. X Gel film, potentially enhance the product stability. It has also been developed for non-ingestible applications such as cosmetic, ostomy pouches, sanitary and healthcare devices. The development and manufacture of XGel films uses a means called “solution casting”.

2) Soluleaves: In this technology, the film is produced in order to release the active ingredients on coming in contact with saliva. This is applied to flavour-release products such as mouth fresheners, confectionery and vitamin products. SOLULEAVES technology can be used to deliver active ingredients to oral cavity efficiently and in a pleasant and easily portable form. The delivery system can be used for the cough/cold, gastrointestinal and pain therapeutic areas as well as nutritional products. SOLULEAVES films can also be designed to adhere to mucous membranes and to release the active ingredients slowly over 15 minutes.

3) Wafertab: WAFERTAB is a drug delivery system that incorporates pharmaceutical actives into ingestible films. It is a patented delivery system that uses a unique process to prepare drug-loaded thin films which can be used in topical or oral application. Active ingredients are incorporated into the film after casting. WAFERTAB system lends itself to many possibilities for innovative drug design, enabling multiple films with different actives to be bonded together.

4) Foamburst: FOAMBURST is a patent granted in September 2004 which is for capsules made of foamed film. Gas is blown into the film during production, resulting in a film with a honeycombed structure. The voids in the film may be gas-filled, empty or filled with other materials to produce specific taste-burst characteristics or to deliver active drugs. The light honeycombed structure results in capsules that dissolve rapidly, causing a melt-in-the mouth sensation. FOAMBURST has attracted from and confectionary manufactures as a mean of carrying and releasing flavours.

5) Micap: Micap signed an option agreement in 2004 to combine its expertise in micro
encapsulation technology with the Bio Progress water-soluble films. The developments aimed at providing new delivery mechanisms for the $1.4bn global market for smoking cessation products (SCPs).

**Pharmacopoeial Status of Oral Films**

Monographs of common dosage forms are provided by the pharmacopoeias (e.g. Ph. Eur., USP). Even though dosage forms for application in the oral cavity such as Medicated chewing gums, Oromucosal preparations, Orodispersible tablets or oral Lyophilisates are included, monographs and specifications for oral films of diverse dissolution kinetics has not yet been established. There are inadequate pharmaceutical technical procedures for analysis in development and quality control of oral films as well. For instance, disintegration and dissolution testing procedures may be provided, but the recommended conditions such as volumes of media do not reflect the natural conditions in the oral cavity.

**Evaluation of Fast Dissolving Films**

1) Organoleptic evaluation
2) Mechanical properties
   a) Thickness
   b) Dry test/tack test
   c) Tensile Strength
   d) Percent Elongation
   e) Tear Resistance
   f) Young’s modulus
   g) Folding endurance
3) Swelling properties
4) Transparency
5) Contact angle
6) Assay/Content uniformity
7) Disintegration time
8) In-vitro Dissolution test

1) **Organoleptic evaluation:** For evaluation of the product, special controlled human taste panels are used. *In vitro* methods of utilizing taste sensors, are being used for this purpose. These *in vitro* taste assessment apparatus and methodologies are well suited for high-throughout taste screening of oral films.

2) **Mechanical properties:**

   **Thickness**
   The thickness of strip can be measured by micrometer screw gauge at different strategic locations. This is essential to ascertain uniformity in the thickness of the film as this is directly related to the accuracy of dose in the strip.

   **Dryness test/tack test**
   Tack is the tenacity with which the strip adheres to an accessory (a piece of paper) that has been pressed into contact with the strip.

   **Tensile Strength**
   Tensile strength is the maximum stress applied to a point at which the strip specimen breaks. It is calculated by the applied load at rupture divided by the cross-sectional area of the strip as given in the equation below:

   \[
   \text{Tensile strength} = \frac{\text{Load at failure} \times 100}{\text{Strip thickness} \times \text{Strip width}}
   \]

   **Percent Elongation**
   When stress is applied, a strip sample stretches and this is referred to as strain. Strain is basically the deformation of strip divided by original dimension of the sample. Generally elongation of strip increases as the plasticizer content increases.

   \[
   \% \text{ Elongation} = \frac{\text{Increase in length of strip}}{\text{Initial length of strip}} \times 100
   \]

   **Tear Resistance**
   Tear resistance of plastic film or sheeting is a complex function of its ultimate resistance to rupture. Basically very low rate of loading 51 mm (2 in)/min is employed to measure the force to initiate tearing. The maximum stress or force (that is generally found near the onset of tearing) required to tear the specimen is recorded as the tear resistance value in Newtons (or pounds-force).

   **Young’s Modulus**
   Young’s modulus or elastic modulus is the measure of stiffness of strip. It is represented as the ratio of applied stress over strain in the region of elastic deformation as follows:

   \[
   \text{Young’s modulus} = \frac{\text{Slope} \times 100}{\text{Strip thickness} \times \text{cross-head speed}}
   \]

   **Folding Endurance**
   Folding endurance is determined by repeated folding of the strip at the same place till the strip breaks. The number of times the film is folded without breaking is computed as the folding endurance value.

3) **Swelling property:** Film swelling study is conducted using simulated saliva solution. Each film sample is weighed and placed in a preweighed stainless steel wire mesh. The mesh containing film sample is submerged into
15ml medium in a plastic container. Increase in the weight of the film is determined at preset time interval until a constant weight is observed. The degree of swelling is calculated using formula:

\[ \alpha = \frac{(wt - wo)}{wo} \]

\( wt \) is weight of film at time \( t \), and \( wo \) is weight of film at time zero.

4) Transparency: The transparency of the films can be determined using a simple UV spectrophotometer. Cut the film samples into rectangles and placed on the internal side of the spectrophotometer cell. Determine the transmittance of films at 600 nm. The transparency of the films can be calculated as follows:

\[ \text{Transparency} = \frac{\log T_{600}}{b} = -\epsilon c \]

Where \( T_{600} \) is the transmittance at 600 nm, \( b \) is the film thickness (nm) and \( c \) is concentration.

5) Contact Angle: Contact angle measurements are performed at the room temperature with a goniometry. A drop of double distilled water was placed on the surface of the dry film. Images of the water droplet were recorded by means of digital camera, digital images are analyzed by the image 1.28v software for angle determination.

6) Assay/ Content uniformity: This is determined by any standard assay method described for the particular API in any of the standard pharmacopoeia. Content uniformity is determined by estimating the API content in individual strip. Limit of content uniformity is 85–115 percent.

7) Disintegration Time: The disintegration time limit of 30 s or less for orally disintegrating tablets described in CDER guidance can be applied to fast dissolving oral strips [44]. Although, no official guidance is available for oral fast disintegrating films/strips, this may be used as a qualitative guideline for quality control test or at development stage. Pharmacopoeial disintegrating test apparatus may be used for this study. Typical disintegration time for strips is 5–30 s.

8) In-vitro Dissolution Test: Dissolution testing can be performed using the standard basket or paddle apparatus described in any of the pharmacopoeia. The dissolution medium will essentially be selected as per the sink conditions and highest dose of the API. Many times dissolution test can be difficult due to tendency of the strips to float on the dissolution medium where paddle system is used.

Storage and Packaging of Films
A variety of storage and packaging options are available for fast dissolving films. The packaging stage provides product flexibility to the drug manufactures. Single packaging is mandatory for films, which are pharmaceutical products; an aluminum pouch is the most commonly used packaging format. APR-Labtec has developed the Rapid card, a proprietary and patented packaging system, which is specially designed for the Rapid films. The rapid card has same size as a credit card and holds three raid films on each side. Every dose can be taken out individually.

Conclusion
Fast Dissolving Films have several advantages over the conventional dosage forms. They are considered as a most important drug delivery system today because of their rapid disintegration, improved dissolution. They combine the greater stability of a solid dosage form and good applicability of the liquid and thus bridge the gap between the two ideas, incorporating positive elements from both solid and liquid dosage forms into an elegant, stable and effective delivery vehicle. So they are of great importance during the emergency cases such as allergic reactions and asthmatic attacks whenever immediate onset of action is desired. So this technology is growing in fast pace challenging most of the pharmaceutical companies to develop oral films for a wide range of active pharmaceutical ingredients.
Table 1: Types of Oral Films and their Properties

<table>
<thead>
<tr>
<th>Property/Sub Type</th>
<th>FlashRelease Water</th>
<th>Mucoadhesive Melt-Away Wafer</th>
<th>Mucoadhesive Sustained Release Wafer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Area (cm²)</td>
<td>2-8</td>
<td>2-7</td>
<td>2-4</td>
</tr>
<tr>
<td>Thickness (µm)</td>
<td>20-70</td>
<td>50-500</td>
<td>50-250</td>
</tr>
<tr>
<td>Structure</td>
<td>Film: single layer</td>
<td>Single or multilayer System</td>
<td>Multi layer system</td>
</tr>
<tr>
<td>Excipients</td>
<td>Soluble, highly hydrophilic polymers</td>
<td>Soluble, hydrophilic Polymers</td>
<td>Low/Non-soluble Polymers</td>
</tr>
<tr>
<td>Drug phase</td>
<td>Solid solution</td>
<td>Solid solution or suspended drug particles</td>
<td>Suspension and/or solid Solution</td>
</tr>
<tr>
<td>Application</td>
<td>Tongue (upper palate)</td>
<td>Gingival or buccal Region</td>
<td>Gingival, (other region in the oral cavity)</td>
</tr>
<tr>
<td>Dissolution</td>
<td>Maximum 60 seconds</td>
<td>Disintegration in a few minutes, forming gel</td>
<td>Maximum 8-10 hours</td>
</tr>
</tbody>
</table>

Table 2: A typical composition contains the following ingredients:

<table>
<thead>
<tr>
<th>AGENTS</th>
<th>CONCENTRATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>DRUG</td>
<td>1-25%</td>
</tr>
<tr>
<td>WATER SOLUBLE POLYMER</td>
<td>40-50%</td>
</tr>
<tr>
<td>PLASTICIZERS</td>
<td>0-20%</td>
</tr>
<tr>
<td>FILLERS, COLOURS, FLAVOURS ETC.</td>
<td>0-40%</td>
</tr>
</tbody>
</table>

Table 3: Compares solvent-casting and HME for the manufacturing of FDFs.

<table>
<thead>
<tr>
<th>PROCESS PARAMETERS</th>
<th>SOLVENT CASTING</th>
<th>HOT MELT EXTRUSION</th>
</tr>
</thead>
<tbody>
<tr>
<td>API selected</td>
<td>Thermoliable, thermostable</td>
<td>Thermostable</td>
</tr>
<tr>
<td>Solvent required</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Process</td>
<td>Hydrous</td>
<td>Anhydrous</td>
</tr>
<tr>
<td>Equipment required</td>
<td>Rollers, coaters</td>
<td>Hot-melt extruder</td>
</tr>
<tr>
<td>Scale-up</td>
<td>May create problems</td>
<td>May not be difficult</td>
</tr>
<tr>
<td>Chance of air entrapment</td>
<td>High chance</td>
<td>Low chance</td>
</tr>
</tbody>
</table>

Table 4: List of marketed products of films

<table>
<thead>
<tr>
<th>Product Category</th>
<th>Ingredients</th>
<th>Applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Appetite suppressant</td>
<td>fucusvesiculosus and guarana extract, gacrciniacambogia</td>
<td>These are top selling natural ingredients associated with weight loss. Cambogia helps to reduce the food intake by suppressing appetite</td>
</tr>
<tr>
<td>Product Name</td>
<td>Description</td>
<td>Use</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------</td>
</tr>
<tr>
<td>Breath freshener strip,</td>
<td>Contain mint flavor and antibacterial agent, cetylpyridinium chloride</td>
<td>It is used as mouth freshener and to stop bad breath</td>
</tr>
<tr>
<td>(Antibacterial strip)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Donezepil Rapid film®</td>
<td>Donepezil Hydrochloride 5 mg and 10 mg.</td>
<td>Treatment of mild to moderately severe dementia of the Alzheimer's type</td>
</tr>
<tr>
<td>Suppress™ Cough strips with menthol</td>
<td>Artificial flavors, ascorbic acid, aspartame, asulfame potassium, carrageenan, diglycerides, fatty acid ester etc.</td>
<td>Temporarily suppresses coughs due to minor throat and bronchial irritation.</td>
</tr>
<tr>
<td>Saliva promoting strips</td>
<td>Fruit acid extracts, range of flavors</td>
<td>It is used in the dry mouth as a side effect of the other medications</td>
</tr>
<tr>
<td>Energy boosters</td>
<td>Caffeine, green tea extract and guarana</td>
<td>The product maintains the energy levels.</td>
</tr>
<tr>
<td>Smoking cessation</td>
<td>Nicotine</td>
<td>To reduce the smoking habit</td>
</tr>
<tr>
<td>Vitamins and food supplements</td>
<td>Various vitamins, minerals and supplements</td>
<td>It is useful for the people who do not like to pop up the tablets or soluble supplements</td>
</tr>
<tr>
<td>Labtec GmbH</td>
<td>Ondansetron 4 mg and 8 mg.</td>
<td>It is used in the prevention of chemotherapy and radiation-induced nausea and vomiting and prevention of postoperative nausea and vomiting</td>
</tr>
<tr>
<td>InnozenInc</td>
<td>Chloraseptic® Relief Strips™</td>
<td>Occasional minor irritation, pain, sore throat and sore mouth</td>
</tr>
<tr>
<td>Chloraseptic® Kids Sore Throat Relief strips</td>
<td>Benzocaine 3 mg, BHT, corn starch, erythritol, FD&amp;C Red 40, hydroxypropyl methylcellulose, malic acid, menthol, monoammonium glycyrrhizinate, cherry flavors, polyethylene oxide, sucralose</td>
<td></td>
</tr>
<tr>
<td>Chloraseptic® Kids Sore Throat Relief strips</td>
<td>Benzocaine 2 mg and menthol, grape flavor, BHT, corn starch, erythritol, FD&amp;C Blue 1, FD&amp;C Red 40, hydroxypropyl methylcellulose, malic acid, menthol, monoammoniumglycyrrhizinate, polyethylene oxide, sucralose</td>
<td>Occasional minor irritation, pain, sore throat and sore mouth</td>
</tr>
</tbody>
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Fig 1: Flow chart of solvent casting method for the preparation of fast dissolving films

Fig 2: Three roll coating unit

Cite this article
References