INTRODUCTION

Drug Delivery Systems (DDS) are a strategic tool for expanding markets/indications, extending product life cycles and generating opportunities. DDS make a significant contribution to global pharmaceutical sales through market segmentation, and are moving rapidly. Drug delivery systems are becoming increasingly sophisticated as pharmaceutical scientists acquire a better understanding of the physicochemical and biochemical parameters pertinent to their performance.

Despite tremendous advancements in drug delivery, the oral route remains the perfect route for the administration of therapeutic agents because of low cost of therapy, ease of administration, accurate dosage, self-medication, pain avoidance, versatility, leading to high levels of patient compliance. Tablets and capsules are the most popular dosage forms.1 But one important drawback of such dosage forms is ‘Dysphagia’ or difficulty in swallowing. This is seen to afflict nearly 35% of the general population and associated with a number of conditions, like parkinsonism, mental disability, motion sickness, unconsciousness, unavailability of water etc. To overcome such problems, certain innovative drug delivery systems, like ‘Mouth Dissolving Tablets’ (MDT) have been developed. These are novel dosage forms which dissolve in saliva within a few seconds, when put on tongue. Such MDTs can be administered anywhere and anytime, without the need of water and are thus quite suitable for children, elderly and mentally disabled patients.

Keywords: Mouth dissolving tablets, MDT.

ABSTRACT

Recent advances in Novel Drug Delivery Systems (NDDS) aim for designing dosage forms, convenient to be manufactured and administered, free of side effects, offering immediate release and enhanced bioavailability, so as to achieve better patient compliance. Though oral drug delivery systems, preferably, tablets are the most widely accepted dosage forms, for being compact, offering uniform dose and painless delivery. Yet, dysphagia is the most common disadvantage of conventional tablets. This is seen to afflict nearly 35% of the general population and associated with a number of conditions, like parkinsonism, mental disability, motion sickness, unconsciousness, unavailability of water etc. To overcome such problems, certain innovative drug delivery systems, like ‘Mouth Dissolving Tablets’ (MDT) have been developed. These are novel dosage forms which dissolve in saliva within a few seconds, when put on tongue. Such MDTs can be administered anywhere and anytime, without the need of water and are thus quite suitable for children, elderly and mentally disabled patients.

Keywords: Mouth dissolving tablets, MDT.

Advantages of MDT

a. No need of water to swallow the tablet.9
b. Can be easily administered to pediatric, elderly and mentally disabled patients.
c. Accurate dosing8 as compared to liquids.
d. Dissolution and absorption of drug is fast, offering rapid onset of action.
e. Bioavailability of drugs is increased8 as some drugs are absorbed from mouth, pharynx and esophagus through saliva passing down into the stomach.11
f. Advantageous over liquid medication in terms of administration as well as transportation
gh. First pass metabolism is reduced, thus offering improved bioavailability and thus reduced dose and side effects.
i. Free of risk of suffocation due to physical obstruction when swallowed, thus
jj. Offering improved safety.
ka. Suitable for sustained/controlled release actives.12
lb. Allows high drug loading.13

MOUTH DISSOLVING TABLETS: A NOVEL APPROACH TO DRUG DELIVERY

TEJVIR KAUR1, BHAWANDEEP GILL2, SANDEEP KUMAR3, G.D. GUPTA3

1Lecturer, Department of Pharmacy, Government Medical College, Patiala, Rayat & Bahra College of Pharmacy, Kharar, HOD Pharmaceutics, Department of Pharmaceutics, ASBASJSM College of Pharmacy, Bela, Ropar Punjab 140111 India Email: tejvironnet@gmail.com, gillbhanwan84@gmail.com, Mr.sandeep1970@rediffmail.com, drgdg@rediffmail.com

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Keywords: Mouth dissolving tablets, MDT.

Mouth dissolving tablet (MDT)

It is a tablet that disintegrates and dissolves rapidly in the saliva, within a few seconds without the need of drinking water or chewing. A mouth dissolving tablet usually dissolves in the oral cavity within 15 s to 3 min. Most of the MDTs include certain super disintegrants and taste masking agents.

Ideal properties of MDT

A Mouth Dissolving Tablet should

a. Not require water or other liquid 1 to swallow.
b. Easily dissolve or disintegrate in saliva within a few seconds.
c. Have a pleasing taste.
d. Leave negligible or no residue in the mouth when administered.
e. Be portable and easy to transport.
f. Be able to be manufactured in a simple conventional manner within low cost.
g. Be less sensitive to environmental conditions like temperature, humidity etc.8,7

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c. Accurate dosing8 as compared to liquids.
d. Dissolution and absorption of drug is fast, offering rapid onset of action.
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lb. Allows high drug loading.13
Main ingredients used in preparation of MDT

Important ingredients that are used in the formulation of MDTs should allow quick release of the drug, resulting in faster dissolution. This includes both the actives and the excipients. Disintegration and solubilization of a directly compressed tablet depend on single or combined effects of disintegrants, water-soluble excipients and effervescent agents.

Excipients balance the properties of the actives in FDDTs. This demands a thorough understanding of the chemistry of these excipients to prevent interaction with the actives. Determining the cost of these ingredients is another issue that needs to be addressed by formulators. The role of excipients is important in the formulation of fast-melting tablets. These inactive food-grade ingredients, when incorporated in the formulation, impart the desired organoleptic properties and product efficacy. Excipients are general and can be used for a broad range of actives, except some actives that require masking agents. Binders keep the composition of these fast-melting tablets together during the compression stage. The right selection of a binder or combination of binders is essential to maintain the integrity and stability of the tablet. The temperature of the excipient should be preferably around 30-35°C for faster melting properties. Further, its incorporation imparts smooth texture and disintegration characteristics to the system. Binders can either be liquid, semi solid, solid or mixtures of varying molecular weights such as polyethylene glycol. The choice of a binder is critical in a fast-dissolving formulation for achieving the desired sensory and melting characteristics, and for the faster release of active ingredients. Commonly available fats such as cocoa butter and hydrogenated vegetable oils can also be used.

The most important ingredients of a mouth dissolving tablets are:

**Super disintegrants**: Use of disintegrants is the basic approach in development of MDTs. Disintegrants play a major role in the disintegration and dissolution of MDT. It is essential to choose a suitable disintegrant, in an optimum concentration so as to ensure quick disintegration and high dissolution rates.14

Super disintegrants provide quick disintegration due to combined effect of swelling and water absorption by the formulation. Due to swelling of superdisintegrants, the wetted surface of the carrier increases, which promotes the wettability and dispersibility of the system, thus enhancing the disintegration and dissolution.15-17 The optimum concentration of the superdisintegrant can be selected according to critical concentration of disintegrant.

Below this concentration, the tablet disintegration time is inversely proportional to the concentration of the superdisintegrant, whereas if concentration of superdisintegrant is above critical concentration, the disintegration time remains almost constant or even increases.18

Sodium starch glycolate, Ac-di-sol(crosscarmellose sodium), Crospovidone, Microcrystalline cellulose, Pregelatinised starch are some of examples of disintegrants.

**Mechanism of action of disintegrants**

The tablet breaks to primary particles by one or more of the mechanisms listed below:-

a. By capillary action
b. By swelling
c. Because of heat of wetting
d. Due to release of gases
e. By enzymatic action
f. Due to disintegrating particle/particle repulsive forces
g. Due to deformation

**a. By capillary action**

Disintegration by capillary action is always the first step. When we put the tablet into suitable aqueous medium, the medium penetrates into the tablet and replaces the air adsorbed on the particles, which weakens the intermolecular bond and breaks the tablet into fine particles. Water uptake by tablet depends upon hydrophilicity of the drug/excipient and on tableting conditions. For these types of disintegrants, maintenance of porous structure and low interfacial tension towards aqueous fluid is necessary which helps in disintegration by creating a hydrophilic network around the drug particles.

**b. By swelling**

Perhaps the most widely accepted general mechanism of action for tablet disintegration is swelling Tablets with high porosity show poor disintegration due to lack of adequate swelling force. On the other hand, sufficient swelling force is exerted in the tablet with low porosity. It is worthwhile to note that if the packing fraction is very high, fluid is unable to penetrate in the tablet and disintegration is again slows down.
Hermann has proposed a particle repulsion theory based on the observation that nonswelling particle also cause disintegration of tablets. The electric repulsive forces between particles are the mechanism of disintegration and water is required for it.

Researchers found that repulsion is secondary to wicking. A tablet. The electric repulsive forces between particles are the mechanism of disintegration and water is required for it. The effervescent blend helps in disintegration of tablet. This explanation, however, is limited to only a few types of disintegrants and can not describe the action of most modern disintegrating agents.

Carbon dioxide released within tablets on wetting due to interaction between bicarbonate and carbonate with citric acid or tartaric acid. The tablet disintegrates due to generation of pressure within the tablet. This effervescent mixture is used when pharmacist needs to formulate very rapidly dissolving tablets or fast disintegrating tablet. As these disintegrants are highly sensitive to small changes in humidity level and temperature, strict control of environment is required during manufacturing of the tablets. The effervescent blend is either added immediately prior to compression or can be added in to two separate fraction of formulation.

Here, enzymes present in the body act as disintegrants. These enzymes destroy the binding action of binder and helps in disintegration.

Actually due to swelling, pressure exerted in the outer direction or radial direction, it causes tablet to burst or the accelerated absorption of water leading to an enormous increase in the volume of granules to promote disintegration.

Another mechanism of disintegration attempts to explain the swelling of tablet made with 'non-swelling' disintegrants. Guyot-Hermann has proposed a particle repulsion theory based on the observation that nonswelling particle also cause disintegration of tablets. The electric repulsive forces between particles are the mechanism of disintegration and water is required for it. Researchers found that repulsion is secondary to wicking.

Hess had proved that during tablet compression, disintegranted particles get deformed and these deformed particles get into their normal structure when they come in contact with aqueous media or water. Occasionally, the swelling capacity of starch was improved when granules were extensively deformed during compression.

This increase in size of the deformed particles produces a break up of the tablet. This may be a mechanism of starch and has only recently begun to be studied.

Sugar based excipients: Sugar based excipients are used for taste masking and as bulking agents. Most of the drugs are having unpleasant or bitter taste. And the basic requirement for designing MDTs is that the drug should not have disagreeable taste. So taste masking is necessary in most of the cases. Sorbitol, mannitol, xylitol, dextrose, fructose, etc. are mainly used. Aqueous solubility and sweetness impart a pleasing mouth feel and good taste masking. But not all sugar-based materials have fast dissolution rate and good compressibility or compactability. However technologies have been developed to make use of the sugar based excipients in the design of fast dissolving tablets.

Other ingredients commonly used are water soluble diluents, lubricants, antistatics, plasticizers, binders, colors and flavors.

**Approaches for preparation of MDT**

Various technologies used in the manufacture of Mouth Dissolving Tablets include:

1. Freeze-drying or lyophilization
2. Sublimation
3. Spray drying
4. Moulding
5. Mass extrusion
6. Direct compression

**Freeze-drying**:

The tablets prepared by freeze-drying or lyophilization are very porous in nature and disintegrate or dissolve rapidly when come in contact with saliva. In this process, water is sublimated from the product after freezing. First of all, the material is frozen to bring it below its eutectic point. Then primary drying is carried out to reduce the moisture to around 4% w/w of dry product. Finally, secondary drying is done to reduce the bound moisture to the required volume. Due to lyophilization, bulking agent and sometimes drug acquire glossy amorphous structure and thus dissolution is enhanced. A tablet that rapidly disintegrates in aqueous solution includes a partially collapsed matrix network that has been vacuum dried above the collapsed temperature of the matrix. The matrix is partially dried below the equilibrium freezing point of the matrix. Vacuum drying the tablet above its collapse temperature, instead of freeze drying below its collapse temperature provides a process for producing tablets with enhanced structural
integrity, while rapidly disintegrating in normal amounts of saliva. However, the use of freeze-drying is limited due to high cost of equipment and processing. Other major disadvantages of the final dosage forms include lack of physical resistance in standard blister packs.

Sublimation: This process involves addition of some inert volatile substances like urea, urethane, naphthalene, camphor, etc. to other excipients and the compression of blend into tablet. Removal of volatile material by sublimation creates pores in tablet structure, due to which tablet dissolves when comes in contact with saliva. Additionally several solvents like cyclohexane, benzene etc. can also be used as pore forming agents. Mouth dissolving tablets with highly porous structure and good mechanical strength have been developed by this method.23,24

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**Diagram:**

**Fig. 1.5: Schematic Diagram of Sublimation Technique for Preparation of MDT**

Spray drying: A highly porous and fine powder is prepared by spray drying an aqueous composition containing support matrix and other components. This is then mixed with active ingredient and compressed into tablet. Allen and Wang25 used this technique to prepare mouth-dissolving tablets, which disintegrated within 20 s.

Moulding: Tablets prepared by this method are solid dispersions. Physical form of drug in the tablets depends on whether and to what extent it dissolves in the wetted mass.26 The drug can exist as discrete particles or micro particles in the matrix. It can dissolve totally to form a solid solution or dissolve partially in the molten carrier and remaining, if any, stays undissolved and dispersed in the matrix.27 Disintegration time, drug dissolution rate and mouth feel will depend on the type of dispersion. Different moulding techniques can be used to prepare mouth-dissolving tablets:

- **Compression moulding:** The powder mixture previously wetted with a solvent like ethanol/water is compressed into mould plates to form a wetted mass.
- **Heat moulding:** A molten matrix in which drug is dissolved or dispersed can be directly moulded into Mouth dissolving tablets.28
- **No vacuum lyophilization:** This process involves evaporation of solvent from a drug solution or suspension at a standard pressure.29

Moulded tablets possess porous structure, which facilitates rapid disintegration and easy dissolution. Moulded tablets offer improved taste due to water-soluble sugars present in dispersion matrix. But moulded tablets lack good mechanical strength and can undergo breakage or erosion during handling and opening of blister packs.20 However, adding sucrose, acacia or polyvinyl pyrrolidone can increase mechanical strength.

**Mass extrusion:** In this technique, a blend of active drug and other ingredients is softened using solvent mixture of water soluble polyethylene glycol, using methanol and then the softened mass is extruded through the extruder or syringe to get a cylinder of product, which is finally cut into even segments with the help of heated blades to get tablets. The dried cylinder can be used to coat the granules of bitter tasting drugs and thereby masking their bitter taste.

**Direct compression:** The disintegrant addition technology23,24 (direct compression) is the most preferred technique to manufacture the tablets due to certain advantages:

- a. High doses can be accommodated and final weight of the tablet can exceed that of other methods.
- b. Easiest way to manufacture the tablets.
- c. Conventional equipment and commonly available excipients are use
- d. A limited no. of processing steps are involved.
- e. Cost-effectiveness.

Tablet size and hardness strongly affect the disintegrant efficacy. Hard and large tablets have more disintegration time than normally required. Very soft and small tablets have low mechanical strength. So, an optimum kind and concentration of disintegrant should be chosen to achieve quick disintegration and high dissolution rates. Above the critical concentration level, however, disintegration time remains approximately constant or even increases.25

**Patented Technologies for preparation of MDT:**

Several technologies are available for preparing Mouth dissolving tablets. But some commercially useful technologies are:

**Zydis technology:** 'Zydis' is the first mouth dissolving dosage form in the market. It is a unique freeze-dried tablet in which the active drug is incorporated in a water-soluble matrix, which is then transformed into blister pockets and freeze dried to remove water by sublimation. Zydis matrix is made up of a number of ingredients in order to obtain different objectives. Polymers such as gelatin, dextran or alginites are added to impart strength during handling. These form a glossy and amorphous structure. Mannitol or sorbitol is added to impart crystallinity, elegance and hardness. Various gums may be added to prevent sedimentation of dispersed drug particles. Water is used as a medium to ensure the formation of a porous dosage form. Collapse protectants like glycine may be used to prevent shrinkage of dosage form during freeze drying and long-term storage.30 If necessary, suspending agents and pH adjusting agents may be used. Preservatives may also be added to prevent microbial growth. Zydus products are packed in blister packs to protect the formulation from environmental moisture. A secondary moisture proof foil punch is often required as this dosage form is very moisture sensitive. When put into the mouth, Zydis unit quickly disintegrates and dissolves in saliva.

**Drawbacks:**

- a. A water insoluble drug can be incorporated only upto 400 mg per tablet or less. On the other hand water soluble drug can be incorporated only upto 60 mg
- b. Fragility and poor stability of dosage form during storage under stressful conditions.

**Orasolv technology:** It is CIMA lab's first mouth dissolving formulation. This technology involves taste masking of active drug. Effervescent disintegrating agent is also used. Conventional blenders and tablet equipments are used for preparation of tablets. Less force of compaction is used for manufacturing so as to obtain soft and quickly disintegrating tablets. There is a limitation of this technology that soft and fragile tablets are formed, therefore needed to be packed in specially designed pick and place package system.

**Durasolv technology:** This too has been developed by CIMA labs. This is one of the suitable technologies to prepare products
requiring low amounts of active drug. This technology uses drug, fillers and a lubricant to prepare the tablet. Conventional tableting equipment is used to prepare the tablet. Due to higher force of compaction used, tablets prepared are rigid. Dosage form can be packaged into conventional packaging system like blisters.

**Wowtab technology:** Yamanouchi pharmaceutical company patented this technology. 'wow' means 'without water'. The active ingredients may constitute upto 50% w/w of the tablet. In this technique, saccharides of both low and high mouldability are used to prepare the granules. Mouldability is the capacity of a compound to be compressed.

Highly mouldable substance has high compressibility and thus shows slow dissolution. The combination of high and low mouldability is used to produce tablets of adequate hardness. Active ingredients are mixed with low mouldability saccharides and then granulated with high mouldability saccharides and then compressed into tablet. The Wowtab product dissolves quickly in 15 s or less. Wowtab product can be packed in both into conventional bottle and blister packs.37

**Flashdose Technology:** This technology is patented by Fujis. This system uses the combination of both Shearform and Ceform technologies in order to mask the bitter taste of the drug. A sugar-based matrix, called 'Floss' is used, which is made up of a combination of excipients (crystalline sugars) alone or in combination with drugs. Nurofen meltlet, a new form of Ibuprofen, combination of excipients (crystalline sugars) alone or in based matrix, called 'Floss' is used, which is made up of a technologies in order to mask the bitter taste of the drug. A sugar

Compressed dosage form Avoid exposure to moisture or humidity. Dissolves in 5 – 45 s depending upon the size of the tablet. Due to higher force of centrifugal force comes into action, which throws the dry drug blend at high speed through small heated openings. Due to the heat provided by carefully controlled temperature, drug blend liquefies to form a sphere, without affecting the drug stability. The microspheres thus formed are compressed into tablets. As the drug and excipients both can be processed simultaneously, it creates a unique microenvironment in which the materials can be incorporated into the microspheres that can alter the characteristics of the drug, such as enhancing solubility and stability.

**NanoCrystal technology**38: For MDI, Elan’s proprietary NanoCrystal technology can enable formulation and improve compound activity and final product characteristics.

Decreasing particle size increases the surface area, which leads to an increase in dissolution rate. This can be accomplished predictably and efficiently using NanoCrystal technology. NanoCrystal particles are small particles of drug substance, typically less than 1000 nanometers (nm) in diameter, which are produced by milling the drug. For fast dissolving tablets, Elan’s proprietary NanoCrystal technology can enable formulation and improve compound activity and final product characteristics. Decreasing particle size increases the surface area, which leads to an increase in dissolution rate. This can be accomplished predictably and efficiently using NanoCrystal technology.

**NanoCrystal™ Fast dissolving technology provides for:**

- Pharmacokinetic benefits of orally administered nanoparticles (<2 microns) in the form of a rapidly disintegrating tablet matrix
- Exceptional durability, enabling use of conventional packaging equipment and formats (i.e., bottles and/or blisters)
- Wide range of doses (up to 200mg of API per unit)
- Employment of non moisture sensitive substances

**Table 1.1: Comparison of fast dissolving techniques**

<table>
<thead>
<tr>
<th>Technology</th>
<th>Novelty</th>
<th>Handling/Storage</th>
<th>Drug release/bioavailability</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZYDIS (R.P. SCHERER, INC.)</td>
<td>First to market</td>
<td>Do not push tablet through foil</td>
<td>Dissolves in 2 - 10 s</td>
</tr>
<tr>
<td></td>
<td>Freeze Dried</td>
<td>Do not use dosage form from damaged package</td>
<td>May allow for pre-gastric absorption</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sensitive to degradation at humidities &gt; 65%</td>
<td>leading to enhanced bioavailability</td>
</tr>
<tr>
<td>ORASOLV (CIMA LABS, INC.)</td>
<td>Unique taste masking</td>
<td>Packaged in patented oil packs</td>
<td>Disintegrates in 5 – 45 s depending upon</td>
</tr>
<tr>
<td></td>
<td>Lightly compressed</td>
<td></td>
<td>the size of the tablet</td>
</tr>
<tr>
<td>DURASOLV (CIMA LABS, INC.)</td>
<td>Similar to Orasolv, but with better mechanical strength</td>
<td>Packaged in foil or bottles</td>
<td>No significant change in drug bioavailability</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Package in bottles</td>
<td>Disintegrates in 5 – 45 s depending upon</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>the size of the tablet</td>
</tr>
<tr>
<td>WOWTAB (YAMANOUCHI PHARMA TECHNOLOGIES, INC.)</td>
<td>Compressed dosage form</td>
<td>Avoid exposure to moisture or humidity.</td>
<td>Disintegrates in 5 – 45 s depending upon</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>the size of the tablet</td>
</tr>
</tbody>
</table>

5
Table 1.2: Some of Promising Drug Candidates for Mouth Dissolving Tablets

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Category</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Antibacterial agents</td>
<td>Giprofloxacin, tetracycline, erythromycin, rifampicin, penicillin, doxycyclin, nalidixic acid, trimethoprim, sulfacetamide, sulfadiazine etc.</td>
</tr>
<tr>
<td>2.</td>
<td>Anthelminetics</td>
<td>Albendazole, mebendazole, thiabendazole, livermectin, praziquantel, pyrantel embonate, dichlorophen etc.</td>
</tr>
<tr>
<td>3.</td>
<td>Antidepressents</td>
<td>Trimipramine maleate, nortriptyline HCl, trazodone HCl, amoxapine, mianserin HCl, etc.</td>
</tr>
<tr>
<td>4.</td>
<td>Antidiabetics</td>
<td>Glibenclamide, glipizide, tolbutamide, tolazamide, gliclizide, chlorpropamide etc.</td>
</tr>
<tr>
<td>5.</td>
<td>Analgesics/anti-inflammatory agents</td>
<td>Diclofenac sodium, ibuprofen, ketoprofen, mefenamic acid, naproxen, oxphenbutazone, indomethacin, piroxicam, phenylbutazone, etc.</td>
</tr>
<tr>
<td>6.</td>
<td>Antihypertensives</td>
<td>Amlodipine, carvedilol, diltiazem, felodipine, nifedipine, prazosin HCl, nimodipine, terazosin HCl etc.</td>
</tr>
<tr>
<td>7.</td>
<td>Antiarrhythmics</td>
<td>Disopyramide, quinidine sulphate, amiodarone HCl, etc.</td>
</tr>
<tr>
<td>8.</td>
<td>Antihistamines</td>
<td>Acetaminophen, caffeine, chlorpheniramine, promethazine, pyrilamine, etc.</td>
</tr>
<tr>
<td>9.</td>
<td>Antiulcer agents</td>
<td>Alprazolam, diazepam, clozapine, amyllobarbitone, inverapam, haloperidol, nitrazepam, midazolam phenobarbitalone, thiouracil, oxazepam, etc.</td>
</tr>
<tr>
<td>10.</td>
<td>Diuretics</td>
<td>Acetazolamide, chlorothiazide, amiloride, furosemide, spironolactone, bunazide, ethanerythric acid, etc.</td>
</tr>
<tr>
<td>11.</td>
<td>Gastro-intestinal agents</td>
<td>Cimetidine, ranitidine HCl, famotidine, domperidone, omeprazole, ondansetron HCl, granisetron HCl, etc.</td>
</tr>
<tr>
<td>12.</td>
<td>Corticosteroids</td>
<td>Betamethasone, beclomethasone, hydrocortisone, prednisolone, prednisolone, methyl prednisolone, etc.</td>
</tr>
<tr>
<td>13.</td>
<td>Antiprotozoal agents</td>
<td>Metronidazole, tinadazole, omdizole, benzidazole, cloquiniol, decoquinate etc.</td>
</tr>
</tbody>
</table>

Future prospects of MDT

Mouth dissolving tablets can offer several biopharmaceutical advantages such as improved efficiency over conventional dosage forms. For example, they require smaller amounts of active ingredient to be effective, improve absorption profiles, and offer better drug bioavailability than regular tablets and capsules. In addition, MDTs may be suitable for the oral delivery of drugs such as protein and peptide-based therapeutics that have limited bioavailability when administered by conventional tablets. These products usually degrade rapidly in the stomach. Because drugs delivered in MDTs may be absorbed in the pregastric sites of highly permeable buccal and mucosal tissues of the oral cavity, they may be suitable for delivering relatively low molecular weight and highly permeable drugs. Future possibilities for improvements in MDTs and drug delivery are bright, but the technology is still relatively new. Several drug delivery technologies that can be leveraged on improving drug therapy from MDTs have yet to be fully realized.

Table 1.3: Marketed Products of MDT

<table>
<thead>
<tr>
<th>Trade Name</th>
<th>Active Drug</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nimulid-MD</td>
<td>Nimesulide</td>
<td>Panacea Biotech, New Delhi, India</td>
</tr>
<tr>
<td>Feldene Fast Melt</td>
<td>Piroxicam</td>
<td>Pfizer Inc., NY, U.S.A</td>
</tr>
<tr>
<td>Zyrof Meltab</td>
<td>Rofecoxib</td>
<td>Zydus, Cadila, India</td>
</tr>
<tr>
<td>Pepcid RPD</td>
<td>Famotidine</td>
<td>Merck and Co., NJ, U.S.A</td>
</tr>
<tr>
<td>Romilast</td>
<td>Montelukast</td>
<td>Ranbaxy Labs Ltd, New Delhi, India</td>
</tr>
<tr>
<td>Torrox MT</td>
<td>Rofecoxib</td>
<td>Torrent Pharmaceuticals, Ahmedabad, India</td>
</tr>
<tr>
<td>Olanex Instab</td>
<td>Olanzapine</td>
<td>Ranbaxy Labs Ltd, New Delhi, India</td>
</tr>
<tr>
<td>Zofran OD</td>
<td>Ondansetron citrate</td>
<td>Glaxo Wellcome, Middlesex, UK</td>
</tr>
<tr>
<td>Mosid-MT</td>
<td>Mosapride</td>
<td>Torrent Pharmaceuticals, Ahmedabad, India</td>
</tr>
<tr>
<td>Febrecol</td>
<td>Paracetamol</td>
<td>Prographarm, Chateauneuf, France</td>
</tr>
<tr>
<td>Macat MLT</td>
<td>Rizatriptan</td>
<td>Merck and Co., NJ, U.S.A</td>
</tr>
<tr>
<td>Zelapar TM</td>
<td>Selegiline</td>
<td>Aminar Corp., London, UK</td>
</tr>
</tbody>
</table>

REFERENCES

40. www.ElanNanoCrystal_Technology.html