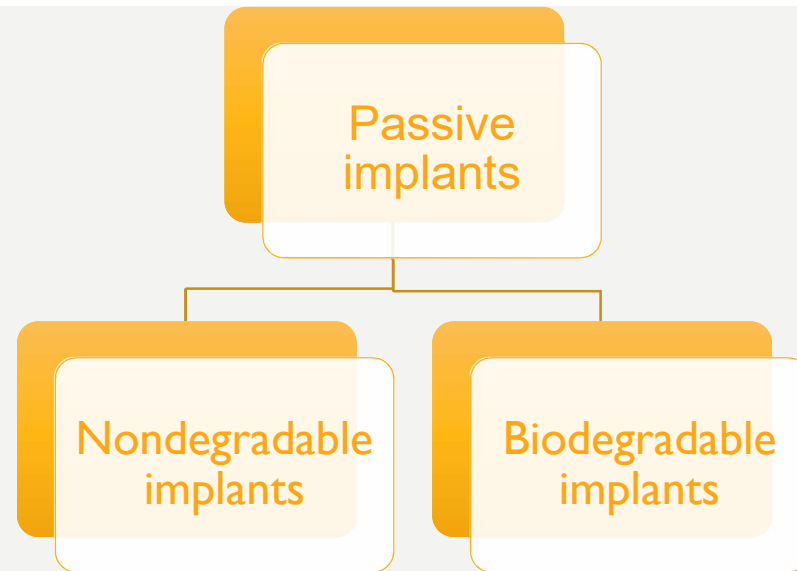


WEEK 11



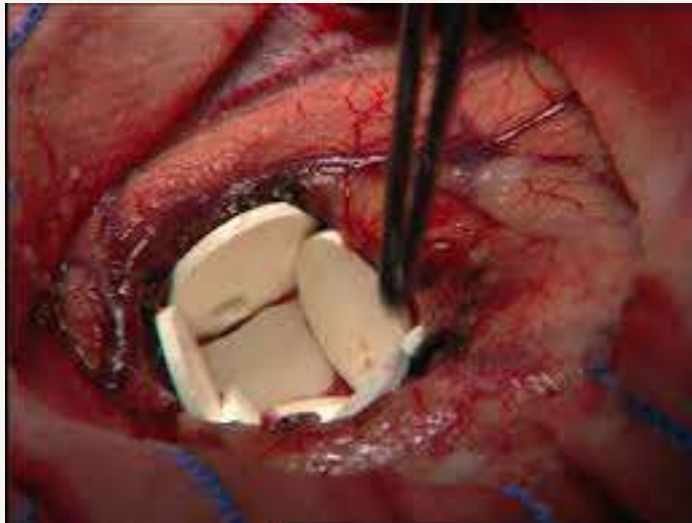
Biodegradable implantable drug delivery systems

biocompatible polymers used for fabricating these delivery systems are eventually broken down into safe metabolites and absorbed or excreted by the body

Polymers include poly(lactic acid) (PLA), poly(lactic-co-glycolic acid) (PLGA), poly(caprolactone) (PCL) or their block copolymer variants with other polymers

Gliadel wafer

- approved by the FDA in 1996
- of biodegradable polyanhydride disks (1.45 cm in diameter and 1.0 mm thick)
- chemotherapeutic drug, carmustine
- biodegradable polyanhydride copolymer



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US Patent Nos. 4,757,128 and 4,789,724
GLIADEL is a registered trademark of GPF Investments, LLC
(polifeprosan 20 with carmustine implant)
Manufactured by Eisai Inc., Research Triangle Park, NC 27709

NDC 62856-177-08


GLIADEL® Wafer

(polifeprosan 20 with carmustine implant)

7.7 mg carmustine/wafer


Each sterile wafer contains 192.3 mg polifeprosan 20 and 7.7 mg carmustine.

Contents:
8 wafers, individually packaged



Store at or below -20°C (-4°F).
Warning: Cytotoxic agent.
See package insert for full prescribing information.
Keep out of the reach of children.

Rx only



GLIADEL® Wafer
(polifeprosan 20 with carmustine implant)
Usual Dose: See package insert.
Handling and Disposal: See package insert.

Lot No:
Exp:

Zoladex

- goserelin acetate
- PLGA or PLA -drug is dispersed in the polymer matrix using hot-melt extrusion method
- prefilled syringe
- biodegradable polyanhydride copolymer
- continuously released over a period of 1 or 3 months



Profact Depot or Suprefact Depot

- buserelin acetate (gonadotropinreleasing hormone agonist)
- PLGA (75:25 molar ratio)
- 2- and 3-month drug release



Passive implants

Dynamic implant systems harness a positive driving force to enable and control drug release. As a result, these are typically able to modulate drug doses and delivery rates much more precisely than passive systems. However, this comes at a higher cost, both in terms of complexity and actual device price.

- Implantable pump systems
- Osmotic pumps
- Propellant infusion pumps

• IMPLANTABLE PUMP SYSTEMS

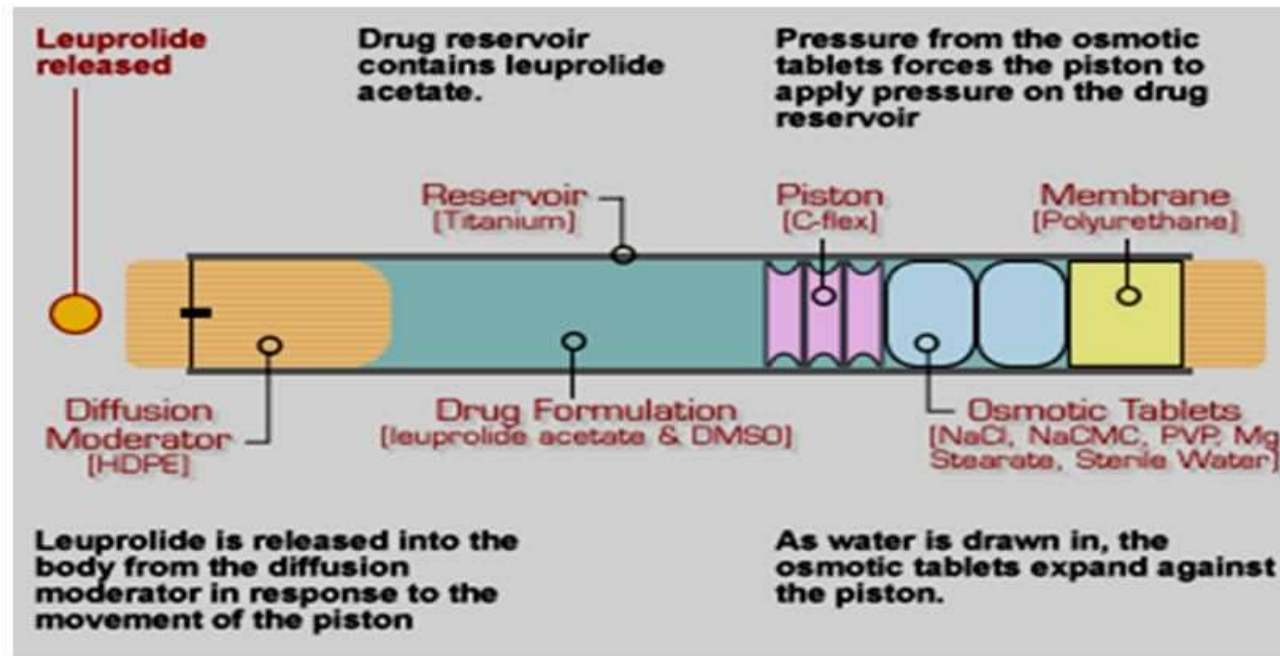
External control of dosing is a requirement for many drugs, a feature that is difficult to obtain when using biodegradable or nondegradable delivery systems. Pump systems have been used to provide the higher precision and remote control needed in these situations. Additionally, they offer a number of advantages, such as evasion of the GI tract, avoidance of repeated injections, and improved release rates (faster than diffusion-limited systems).

Implantable pumps primarily utilize **osmosis, propellant-driven fluids, or electromechanical** drives to generate pressure gradients and enable controlled drug release.

Osmotic pumps

The design comprises a drug reservoir surrounded by a semipermeable membrane, which allows a steady inflow of surrounding fluids into the reservoir through osmosis. A steady efflux of the drug then ensues via the drug portal, an opening in the membrane, as a result of the hydrostatic pressure built on the drug reservoir. Nearly constant or zero-order drug release is maintained until complete depletion of the drug packaged in the reservoir

Viadur-DUROS Device

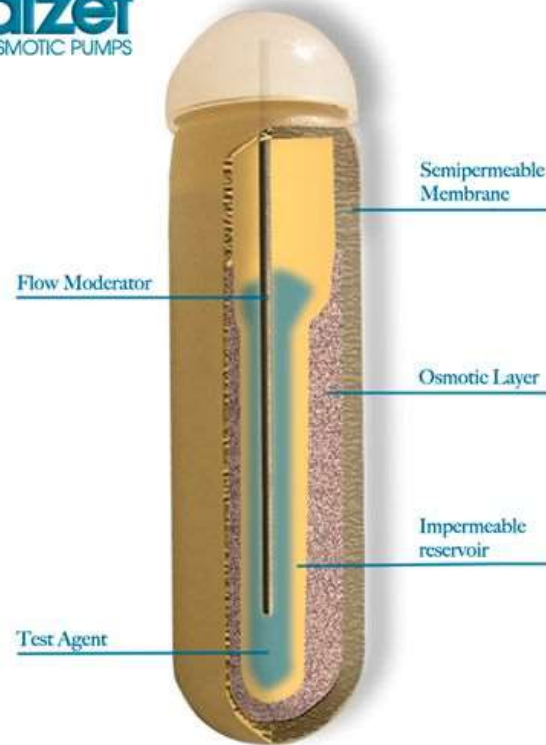


Viadur implant delivers leuprolide for the treatment of prostate cancer

The system is under further development for the delivery of exenatide and has also been investigated for the delivery of other drugs, including interferon (OMEGA DUROS system), sufentanil (Chronogesic system), and other opioids.

Principle of Operation

alzet
OSMOTIC PUMPS



www.alzet.com

ALZET Osmotic Pumps

ALZET pumps have 3 concentric layers:

- Rate-controlling, semi-permeable membrane
- Osmotic layer
- Impermeable drug reservoir

ALZET pumps work by osmotic displacement. Water enters the pump across the outer, semi-permeable membrane due to the presence of a high concentration of sodium chloride in the osmotic chamber. The entry of water causes the osmotic chamber to expand, thereby compressing the flexible reservoir and delivering the drug solution through the delivery portal.

www.alzet.com


DURECT

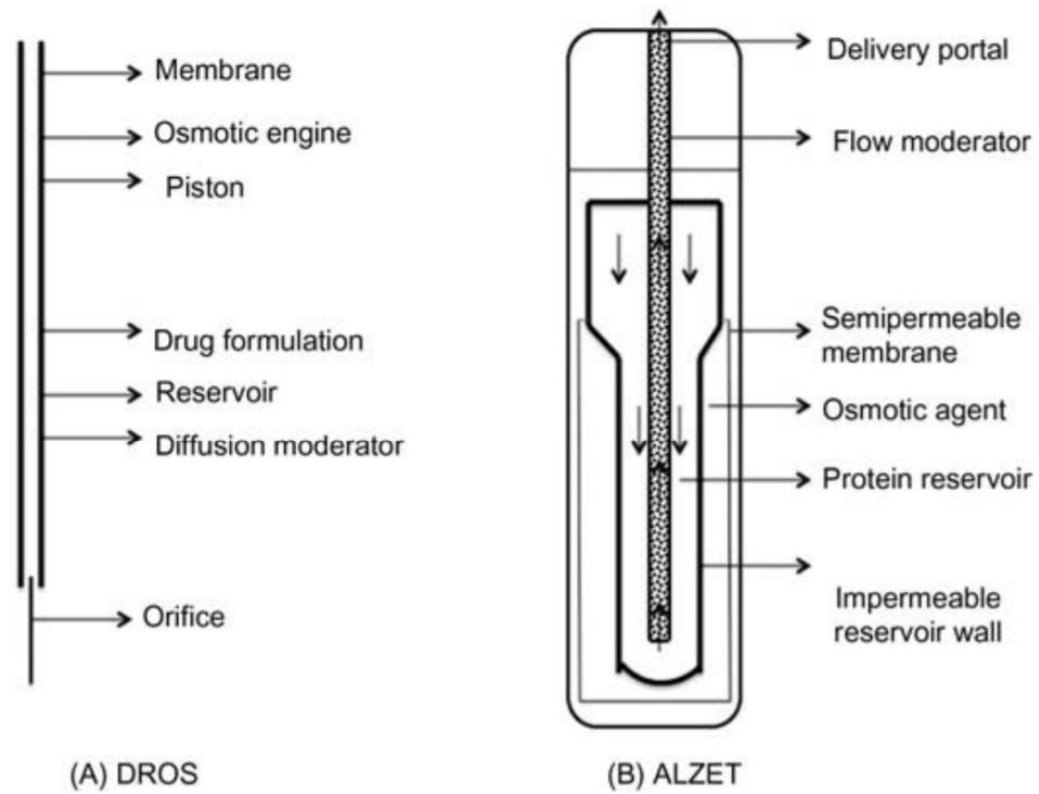


FIGURE 13.2

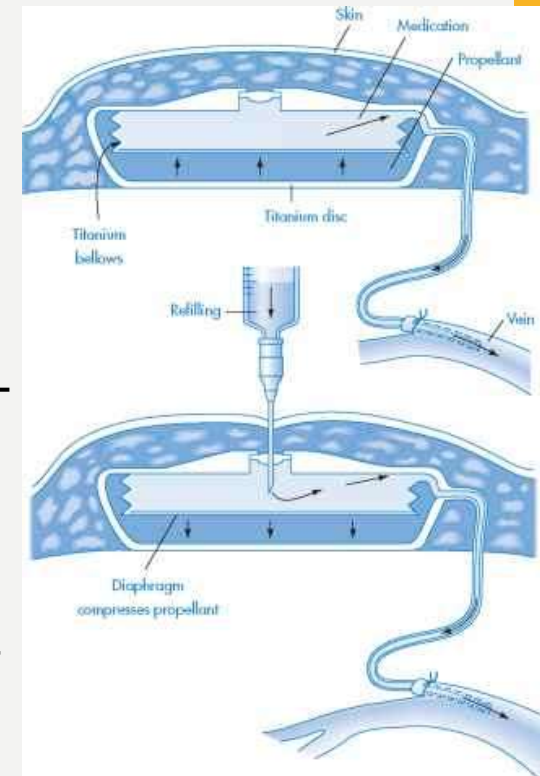
Schematic of osmotic pumps for drug delivery (A) DUROS and (B) ALZET.

Propellant infusion pumps

- The volume of drug that they can release limits the usage of osmotic pumps
- Propellant gas is used instead of an osmotic agent

Infusaid

- Device consists of a small titanium disc which is divided into two chambers by cylindrical titanium bellows that form a flexible but impermeable barrier between the compartments.
- The **outer** compartment contains Freon (chlorofluorocarbon propellant);
- The **inner** compartment contains the infusate and connects via a catheter to a vein or artery through a series of filters and flow-regulating resistant elements.
- The vapour pressure above the liquid propellant remains constant because of the relatively constant temperature of the body, and hence a constant pressure is exerted on the bellows, ensuring a constant rate of delivery of infusate into the bloodstream.
- utilized for insulin delivery, anticoagulant therapy, and cancer chemotherapy

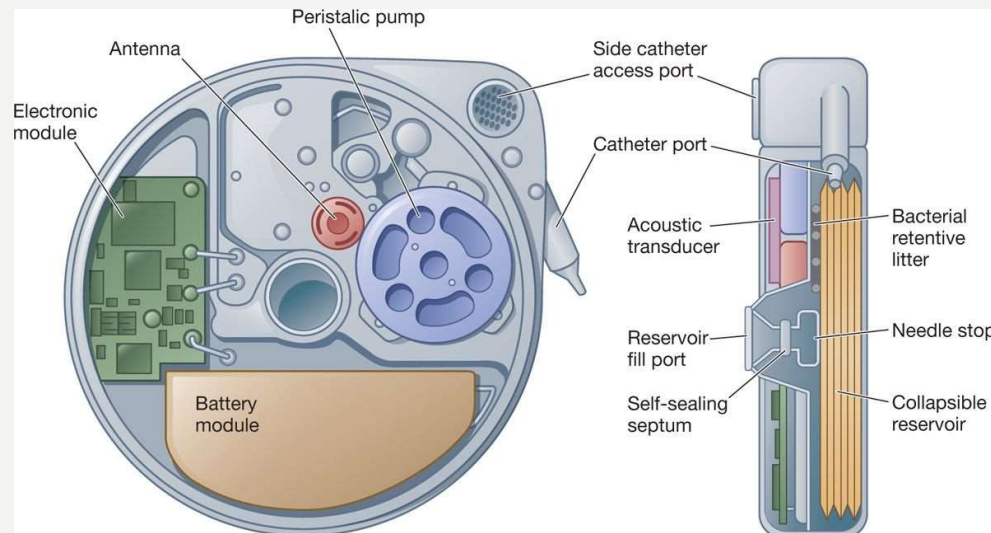




Electromechanical Systems

While osmotic and propellant-driven constant pressure pumps work well for small volumes of medication, this may be a severe limitation for certain chronic diseases requiring daily infusion of medication, precluding their use over long timespans. In such cases, it may be necessary to consider larger implants, wherein the storage capacity of the pump may be replenished from time to time, while the pumping mechanisms stay implanted. By necessity, this implies the use of electrically powered mechanical pumps, typically with moving parts and advanced control systems

Synchromed

- a peristaltic pump implant
- external micro-electronic control of the delivery rate
- pain management using intrathecal delivery of opioids
- The pump consists of an outer titanium shell that encases the pump mechanism and controller, a reservoir holding the drug solution, and a battery.
- It can be conveniently refilled with a needle and syringe via a silicone rubber septum on the system.
- The system is typically implanted in the abdominal cavity.





Micro electromechanical systems (MEMS) technology enables the manufacture of small devices using microfabrication techniques, similar to that used to fabricate silicon-based computer chips. MEMS technology has been used to construct micro-reservoirs, micropumps, nano-porous membranes, nanoparticles, valves, sensors, micro-catheters, and other structures using biocompatible materials appropriate for drug administration.