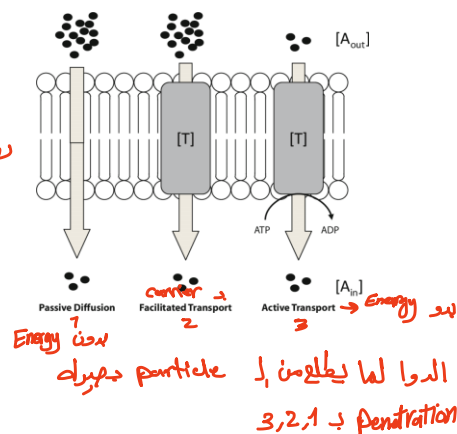


Introduction to nanoparticle properties

Biological transport of NPs

- For drug delivery, most of the sites are accessible through either microcirculation by blood capillaries or pores present at various surfaces and membranes. *microcirculation* *لنقل الدواء*
- Most of the apertures, openings, and gates at cellular or subcellular levels are of nanometer size; hence, NPs are the most suited to reach the subcellular level. *nanometer*
- One of the prime requirements of any delivery system is its ability to move around freely in available avenues and by crossing various barriers that may come in the way.



15

15

لهم اعرف بال penetration ملقحة دخول لا cell
وتحتم يمكن يدخل مع اد nano كلها ويرض لا cell

Table 9 Approximate Sizes of Components in a Typical 20- μ m Human Tissue Cell

Component	قرآن	Size (nm)
Ribosomes		25
Golgi vesicles		30-80
Secretary vesicles		100-1000
Glycogen granules		10-40
Lipid droplets		200-5000
Vaults		55
Lysosomes		500-1000
Proteasomes		11
Peroxisomes		500-1000
Mitochondria		500-1000
Superfine filaments		2-4
Microfilaments		5-7
Thick filaments		15
Microtubules		25
Centrioles		150
Nuclear pores		70-90
Nucleosomes		10
Chromatin		1.9

للمقارنة مع الحفظ

اذا بي افكر بواستي يدخل ←
للنفاذ للنقل اعرف انهم pores
تاعهم 70-90

16

16

حسب ال target tissue لحدود ال size لا nano

7

Solid NPs

- Solid NPs are solid constructs in the nanometer range, and can be prepared by a number of different manufacturing methods which generally involve either:

18 راحة سلاية
 - size reduction of particles (e.g. by milling) to within the nanoparticle range,

- commonly used to prepare drug particles in the nanosize range where there is no carrier material added فقط Drug بدون carrier او Polymer

- molecular agglomeration (e.g. by precipitation methods) to form NPs

- more commonly used to prepare nanoparticle carriers in which drug is loaded.



* اكبر / ابلت بال molecular level وامله agglomeration من precipitation يعني
 يعني solution و ابلت بال precipitation بحيث اتحكم ب size crystal

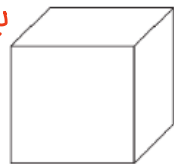
17
 * يمكن احضر solution في drug و polymer ذايين بعد ان اعمل precipitation فيكون في nano carrier

صورة بتوضح كيف اروح عال NPs

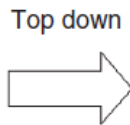
Solid NPs

Top down:

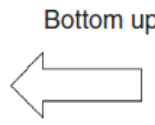
يعني ابلت باستي كبير
 بعمله milling لصاوهل
 لا nano



Bulk materials



Nanomaterials



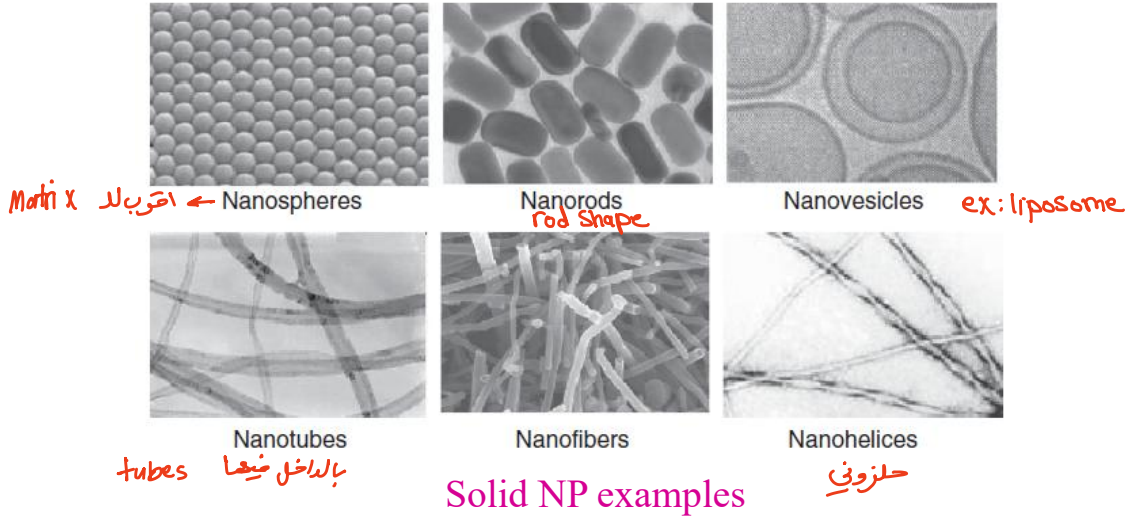
Atoms or molecules

Bottom up: ابلت باستي اصغر

من في nano رايه atomic or molecular level
 من ملحه و اكبره لصاوهل
 - nano

Two basic methods to manufacture nanomaterials.

Solid NPs



19

19

مثال عام Top Down

NPs by wet ball milling

- The preparation of pharmaceutical NPs by **wet (bead, or media) milling** reduces drug particles to a mean particle size less than $0.4 \mu\text{m}$, more commonly 100 or 200 nm.
- In this process, drug is wet milled as a suspension in **water**, or in a medium such as **safflower oil**, **ethanol**, **t-butanol**, **hexane**.
- The milling dispersion also contains one or more surface modifiers which adsorb onto the freshly formed surfaces of the drug and prevent agglomeration through **steric** and/or **electrostatic** stabilization.

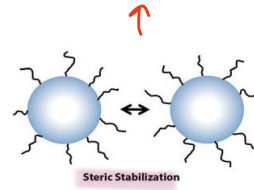
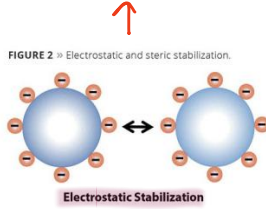
20

20 * *Drug milling مع water او 1, 2, 3, 4 بنقل milling ب ball mill واحياناً يمكن*

steric/electrostatic stabilization. surface modifiers to adsorb onto surface. يمنع الagglomeration لا تغا بغير

في Charges بتعمل repulsion

وجود chains بتخلع لجدران يحزن



Aspect	Electrostatic Stabilization	Steric Stabilization
Mechanism	Repulsion from surface charges plus electrical double layer	Physical barrier from adsorbed polymers/surfactants
Key Force	Coulombic (electrostatic)	Entropic repulsion: Overlap restricts polymer chain conformations Osmotic repulsion: Solvent molecules rush in to dilute it, pushing particles apart.
Common Stabilizers	Ionic surfactants, pH adjustment <i>charge احفظ على, buffer</i>	Non-ionic polymers (e.g., PEG, PVP)
Advantages	Simple, effective in low-salt water	Robust in high-salt or non-aqueous media <i>ما له علاقة بال pH</i>
Disadvantages	Sensitive to electrolytes / Ionic strength <i>sensitive to</i>	Requires thick polymer layer; higher cost
Best For	Aqueous systems with low ionic strength <i>charge وجودك Ion. تهرب ال charge aggregation</i>	High-salt, organic solvents, or broad pH <i>ما بس ال تركيز بال ال pH ال system ما له علاقة ال Ions</i>

21

* Electrical double layer: nanoparticle surface layer في suspension في layer ال particle بتحيط صفا قبل ال
 * osmotic repulsion: movement of solvent from low conce to high concentration
 = (solute)
 = (nano)

NPs by wet ball milling

- Typical surface modifiers include low viscosity *pluronic, PVPs, HPC polymer, hydroxypropyl celluloses, HPMC polymer, hydroxypropyl methylcelluloses, PEGs, lecithin, dextran, aerosol OT, Tween 80, sodium lauryl sulfate, docusate sodium, and (sodium deoxycholate) bile salt*
Aerosol للذرات ال surface activity
- Drug concentrations as high as 40% can be milled, and milling time can range from minutes to days depending on the energy put into the milling process.
milling time حسب ال energy يكون ال
- Milling can be done under refrigerated conditions which minimize thermal degradation.
** ال ال milling لتباج ال Cool jacket ال refrigerator له ال يهرب في Heat*

22

22

يمكن ال صيف surface modifiers ال low viscosity ال عن ال ال aggregation

- Aerosol OT (AOT), or sodium bis(2-ethylhexyl) sulfosuccinate, is a highly effective anionic surfactant

22

Slide 18 هاد اول نوع من NP اسم nano crystal
Top down
Bottom up

The mill can be operated in a batch or a recirculation mode.

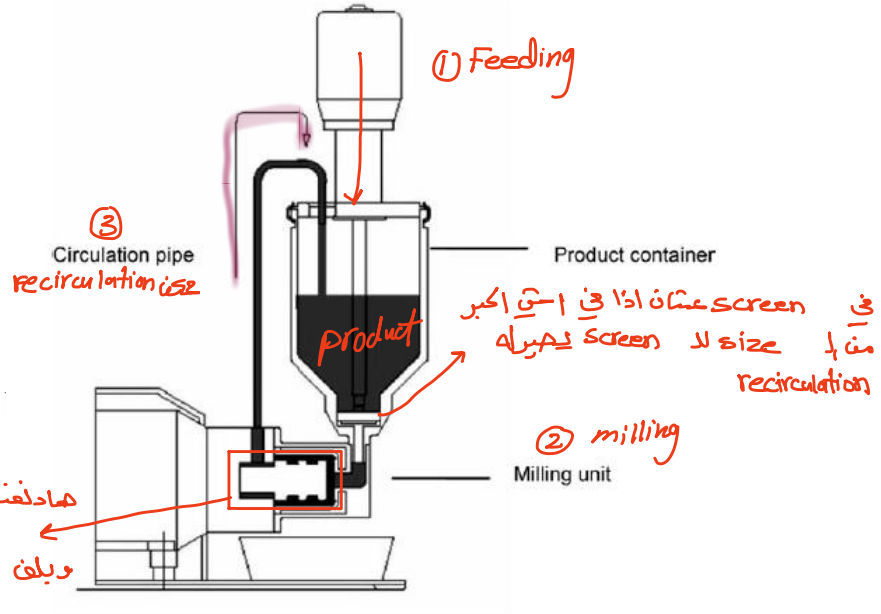
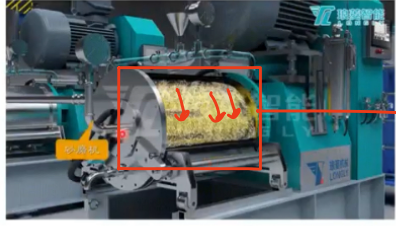


Figure 1 DISPERMAT® SL: schematic view of a bead mill using recirculation method. Source: From Ref. 21.

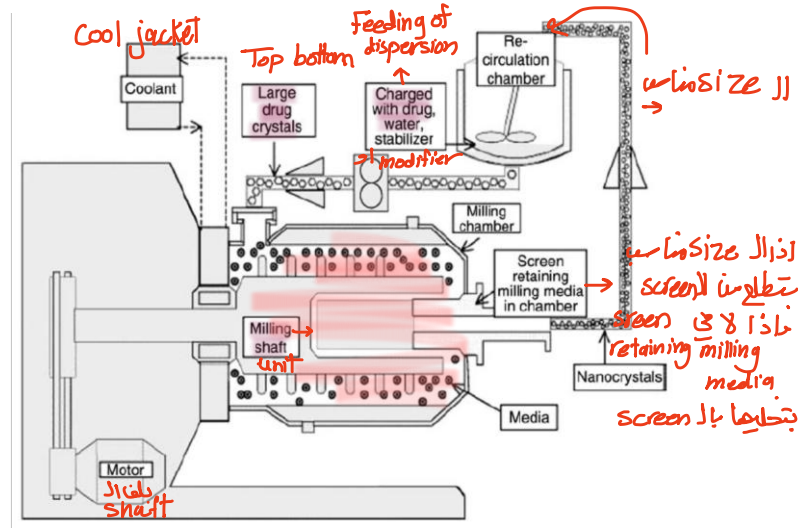
ال mill اسما bead mill

الجزء الي عليه هاد بلف و فيه beads في attrition عالي

كانت تحيي عن Top down ← Top Bottom

The media milling process

- The milling chamber charged with polymeric media is the active component of the mill.
 - A crude slurry consisting of drug, water, and stabilizer is fed into the milling chamber and processed into a nanocrystalline dispersion.
 - The typical residence time required to generate a nanometersized dispersion with a mean diameter <200 nm is 30–60 min or days
- Drug + stabilizer
Drug + water + stabilizer
nanocryst
water
Freeze drying
micro filtration



(From Liversidge, E.M.; Liversidge, G.G.; Cooper, E.R. Eur. J. Pharm. Sci. 2003, 18, 113–120).

هاد نفسها bead mill

منش للوصف ليس حكت الي مكتوب والي مصدر

Table 1. Marketed products containing drug nanocrystals.

Trade name	INN name	FDA approval	Nanosizing technology	Company	Administration route
Rapamune®	Sirolimus	2000	NanoCrystal® (WBM)	Pfizer (Wyeth)	Oral
Emend®	Aprepitant	2003	NanoCrystal® (WBM)	Merck	Oral
Tricor® Lyphanyl®	Fenofibrate	2004	NanoCrystal® (WBM)	Fournier Pharma, Abbott Laboratories	Oral
Triglide®	Fenofibrate	2005	IDD-P® (HPH) high- pressure homogenization	Sciele, Shionogi Pharma Inc.	Oral
Megace® ES	Megestrol acetate	2005	NanoCrystal® (WBM)	PAR Pharmaceuticals	Oral
Invega® Sustenna® Xeplion®	Paliperidone palmitate	2009	NanoCrystal® (WBM)	Janssen	Parenteral, Intramuscular

25

①

للزمن تعرف ان هذا product أسسه nano crystal

25

- ② ممكن بدل ال ball mill او bead mill استخدم high- pressure homogenizer يعني nano crystal
- ③ oral او IM ما بغير IV عن Dispersion
- ④ غالباً الادوية ال solubility العاقلية

Solid polymeric NPs

- Solid NPs can be formed from polymers with the drug incorporated within the polymer matrix or associated onto the particle surface.
- Polymeric NPs are generally formulated from natural or synthetic polymers with the most commonly studied polymers being those which are **biodegradable**, such as:
 - poly(lactide-co-glycolide) (PLGA),
 - polylactic acid (PLA),
 - polycaprolactone (PCL)
 - chitosan

الذبح ياهل الي افدناه Nanocrystal هما الذبح التي Polymeric nano par

26

26

لهن ال polymer جزء رئيسي بار nano بلتة فتح بسلايد 54

Solid polymeric NPs

Several nanoencapsulation methods that have been adapted to pharmaceutical use.

Example: Emulsion evaporation method

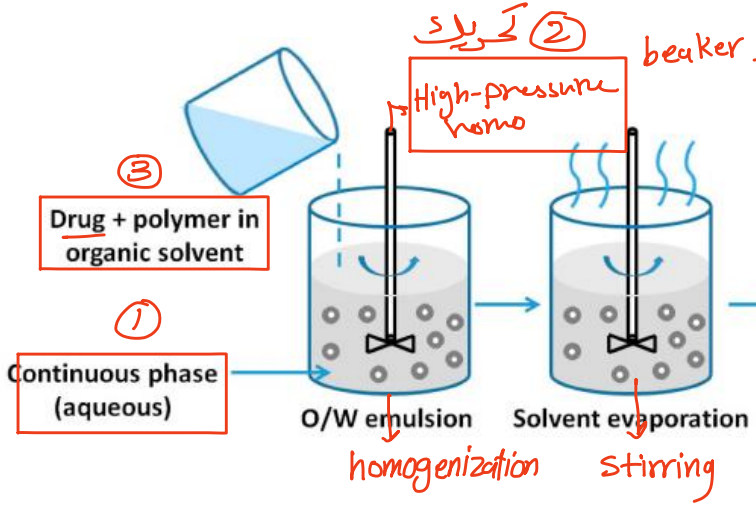
أسهل طريقة لتحضير الـ polymeric NPs

قوائم

- Drug and polymer are dissolved in a volatile organic solvent.
- The mixture is emulsified in an aqueous solution to yield very small droplets size (e.g. by high pressure homogenization). *Emulsifier: tween 80, SLS, Docusate*
- The resulting emulsion is stirred until most of the organic solvent evaporates, leaving solid NPs that may be washed with water and freeze-dried.
- To facilitate solvent evaporation, the emulsion is often heated slightly above the boiling point of the solvent. For example, when methylene chloride (boiling point: 39.8 °C) is used as an organic solvent, the emulsion is heated to ~40 °C.

3) الـ Drug يا ذائب بال organic phase او aqueous phase و بنجيب Beaker بنجيب فيه مثلاً
 27 [Homogenization] High-pressure homoge. و بنجيب فيه مثلاً
 و بنجيب فيه مثلاً aqueous phase و بنجيب فيه مثلاً organic phase

لجدين بيبلت ابيض الـ Drug والـ organic و يجانس و يعطى Emulsion بيبلت اعل solvent evaporation
 و يجانس و يعطى homogenous و لما يصير homogenous



Emulsion evaporation method

عن طريق الـ stirring بنجيب الـ beaker
 على stirrer على حرارة 35-40
 الـ solvent مثلاً Dichloromethane
 فلذلك اقرير به solvent evaporation و nano solidification
 و بنجيب فيه مثلاً aqueous phase
 و بنجيب فيه مثلاً water
 بعد microfiltration و بنجيب فيه مثلاً washing
 و بنجيب فيه مثلاً Room Temp Drying
 و بنجيب فيه مثلاً micro size

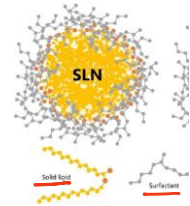
اذا كان الـ size بالـ nano و بنجيب فيه مثلاً evaporation و بنجيب فيه مثلاً Freezing و بنجيب فيه مثلاً (lyophilization) Freeze-drying

28
 → Emulsion: 2 phases (oil + water) + Emulsifying agent
 * Nano emulsion evapor. - عن ادخرا الـ Emulsion العادي
 لازم القرير يكون سويح جداً مثلاً الـ high-pressure homogenizer سويحته بنجيب فيه 15000-20000

في نوع بدل ال polymeric NP في lipid NPs

Solid lipid NPs

- These are NPs made from **solid (high melting point) lipids** dispersed in an aqueous phase.
- Examples of lipids used include:
 - solid triglycerides,
 - saturated phospholipids
 - fatty acids
- The drug is incorporated within the lipid matrix of the particle or by attaching the drug to the lipid nanoparticle surface.
- Solid lipid nanoparticle dispersions have been developed for **parenteral, oral, ocular, dermal** and **cosmetic** applications.
- They can be prepared on a large-scale by homogenization to disperse the lipid into an aqueous environment.



يمكن يكون
بإظلم Surfactant

يمكن اردوا بالداخل او
covalent crosslink on surface
اي على 12

29

ملفوفة الحفيرة : تقين الحفيرة by homogenization / في lipid phase و aqueous phase و surfactant

بحفيرة هم ب large scale homo ب Aqueous Environment

Protein NPs

protein carrier ال

- The first commercial product based on protein nanotechnology was **Abraxane[®]** (paclitaxel).
- Abraxane[®]** consists of 130 nm particles of **albumin-bound paclitaxel**.
- The drug, paclitaxel, has a (low water solubility) and requires addition of solubility-enhancing agents to allow its clinical use.
- The albumin functions to coat the paclitaxel and provide colloidal stabilization to the drug and avoid the toxicity issues associated with the use of solubilizers.

في حالة ال Albumin بحفيرة ال Colloidal suspe ب Albumin بعه Stabilization
فما سيجم Solubilizer

30

30

nab™-paclitaxel complex
130 nm in size¹⁰

nab™-paclitaxel individual molecule
4-14 nm in size^{6,11}

Albumin large protein

Paclitaxel → dispersion of paclitaxel

A single molecule of albumin can bind up to 6 or 7 molecules of paclitaxel*

Nab-Paclitaxel (Abraxane)

31

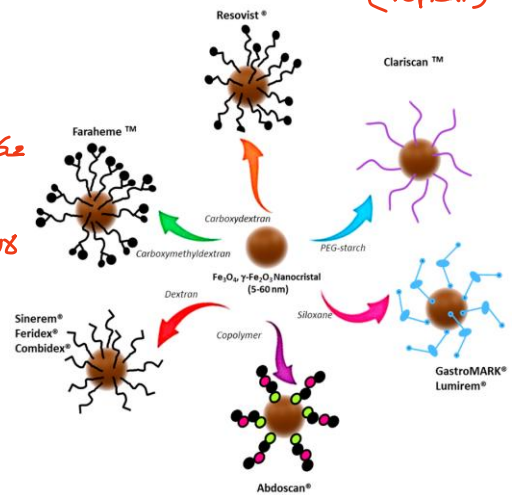
organic → lipid NP
→ polymeric
→ protein

31

Inorganic NPs

- NPs can be fabricated from inorganic materials including metal oxides, metal sulphides, carbon nanotubes, calcium phosphate and ceramics.
- Disadvantage: they are not biodegradable and so have a more limited application.
- Example: **Abdoscan®** is an iron oxide nanoparticle formulation which is administered orally and can be used for magnetic resonance imaging (MRI) diagnostics of the bowel, as it is a superparamagnetic.

في علاج كثير ابحاث
* silver NP: wound healing (topical)



32

32

* Inorganic Fe²⁺ metal زي Ferrous او الي فوق

* Abdoscan النوع الوحيد الموجود بعبارة عن Ferrous و ال Ferrous له magnetic properties
وهي الخاصة بشفه منها بالتصوير الرنيني مثلًا

36 **Polymer-drug conjugates** / اقتران بوليمر

قريب علاج
Pro drug

□ To improve the drug solubility and/ or the delivery of drugs, drug molecules may be conjugated to polymers producing **polymer-drug conjugates**.
↓
لخصها linkage مع polymer

□ These polymer-drug conjugates are considered as new chemical entities in their own right and, as their overall size is generally below 100 nm, these systems can be classified within the general area of nanotechnology.
لأنه صحت دوا لحاله
Drug + polymer

33

33

Polymer-drug conjugates

A polymer-drug conjugate can be described as being built of **three basic components**:

1) **A water soluble polymer backbone.**

Examples: PEG, poly(ethyleneimine) (PEI), PVP, Polyvinylalcohol (PVA), poly(glutamic acid) (PGA) hydroxypropylmethacrylate (HPMA) copolymers.

2) **A linker group.** بتميز ال polymer بأكثر بديل ال Drug

- to help avoid the therapeutic action of the drug being blocked by the polymer
- can also be designed to be cleaved under certain conditions, such as changes in pH, enzymatic degradation or hydrolysis.

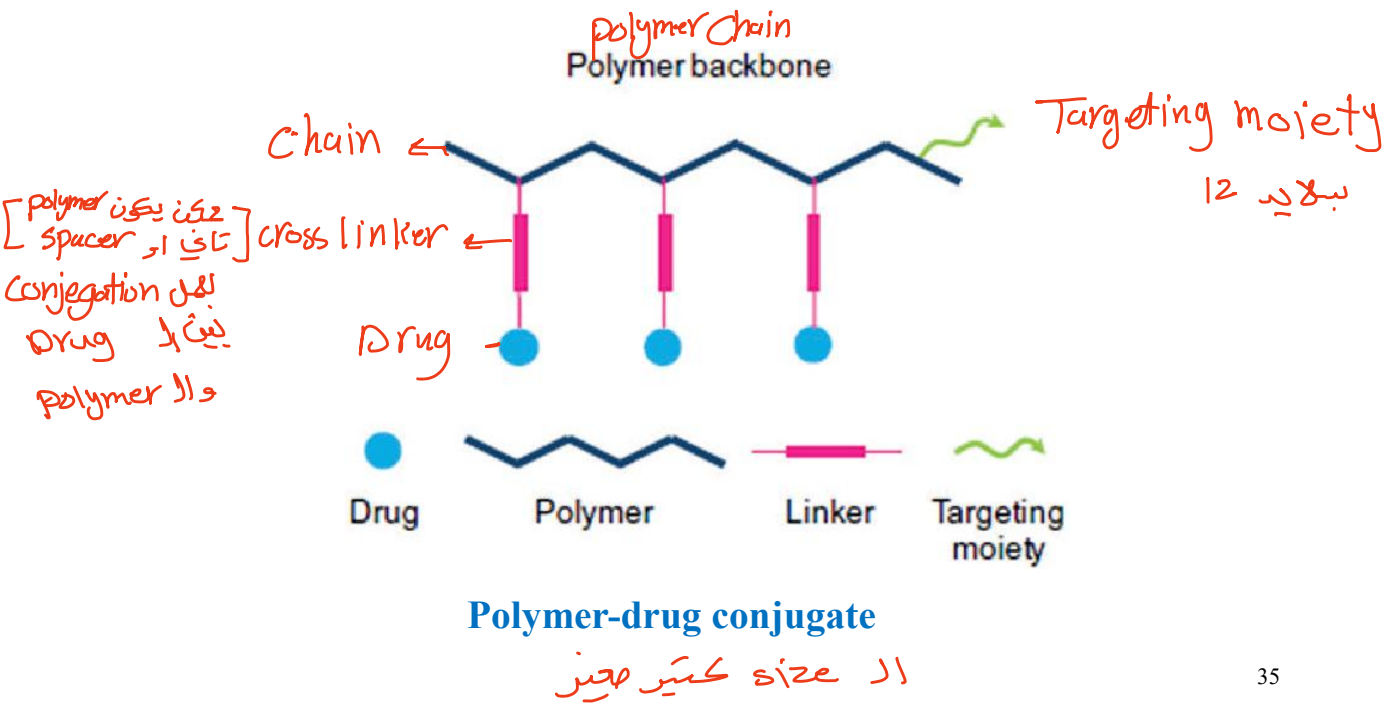
ممكن ال polymer ليعمل interaction مع ال
linker ف receptor يمنع حاد يصير

3) **Drug**

34

34

* linker ممكن يتفعل يكون ل condition معينة مثلاً يصير cleavage عند pH معينة او
وجود Enzyme معين او يصير hydrolysis اذا كان Ester or amid linkage



35

35

Polymer-drug conjugates

قرآنهم

Water soluble polymer backbone

Examples:

- **Synthetic:** poly(ethyleneglycol) (PEG),
- poly(ethyleneimine) (PEI),
- Poly (vinylpyrrolidone) (PVP),
- Polyvinylalcohol (PVA),
- poly(glutamic acid) (PGA)
- hydroxypropylmethacrylate (HPMA) copolymers.
- **Natural** : dextran, chitosans, hyaluronic acid and proteins can be used.

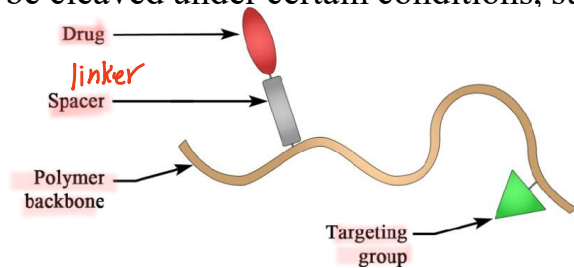
36

36

Polymer-drug conjugates

A linker group

- Whilst a drug can be directly covalently bonded to a polymer, it is more common to attach the drug via a *linker* or *spacer group*, to help avoid the therapeutic action of the drug being blocked by the polymer.
- The linker can also be designed to be cleaved under certain conditions, such as:
 1. changes in pH,
 2. enzymatic degradation or
 3. hydrolysis.



37

37

Polymer-drug conjugates

A linker group

- This property can be used to promote the triggered release of the drug from the polymer conjugate under suitable conditions, thereby enhancing drug targeting. *حسب pH او وظيفه Enzyme*
- Examples of linker groups that can be used include: *amidase ← amid*
 - amine *كلم بصير*
 - carbamate
 - ester groups, *esterase ← ester*
 - amide (the most common option)

كلم بصير
water
hydrolysis

38

38

عادةً لا يتم تصنيع الأدوية عادةً ، يكون لـ Cancer أو CNS نهائياً ، باركسون

Polymer-drug conjugates

Drug

موجودين بالورد

- Anti-cancer chemotherapeutic agents e.g. doxorubicin and paclitaxel
 - polymer conjugates can improve delivery and reduce unwanted side effects or these drugs which have narrow therapeutic windows.
- Proteins e.g. L-asparaginase or interferons.
 - By conjugating proteins to polymers it is possible to increase their half-life by protecting the proteins from enzyme degradation and reducing clearance rates.

* NP لا يقل الـ E.D لأنه مخيمه امدوا قليلة جداً وكلما كان targeted يقل الـ toxicity

39

* proteins يمكن بصيرها Enzyme Degradation فلما نعمل linkage لصي الـ proteins
polymer بغيرها protection من الـ Degradation

Liposomes and bilayer vesicles

أصحة على 42

- Liposomes are closed spherical vesicles consisting of an aqueous core surrounded by one or more bilayer membranes (lamellae) alternating with aqueous compartments.
- These bilayer membranes can be composed of **natural** or **synthetic** amphiphilic lipids, and commonly **phospholipids** are employed for the formulation of liposomes, however a range of amphiphilic lipids can be used.

Triglycerides أو Cholesterol أو Lecithin يمكن استعمالهم

40

40

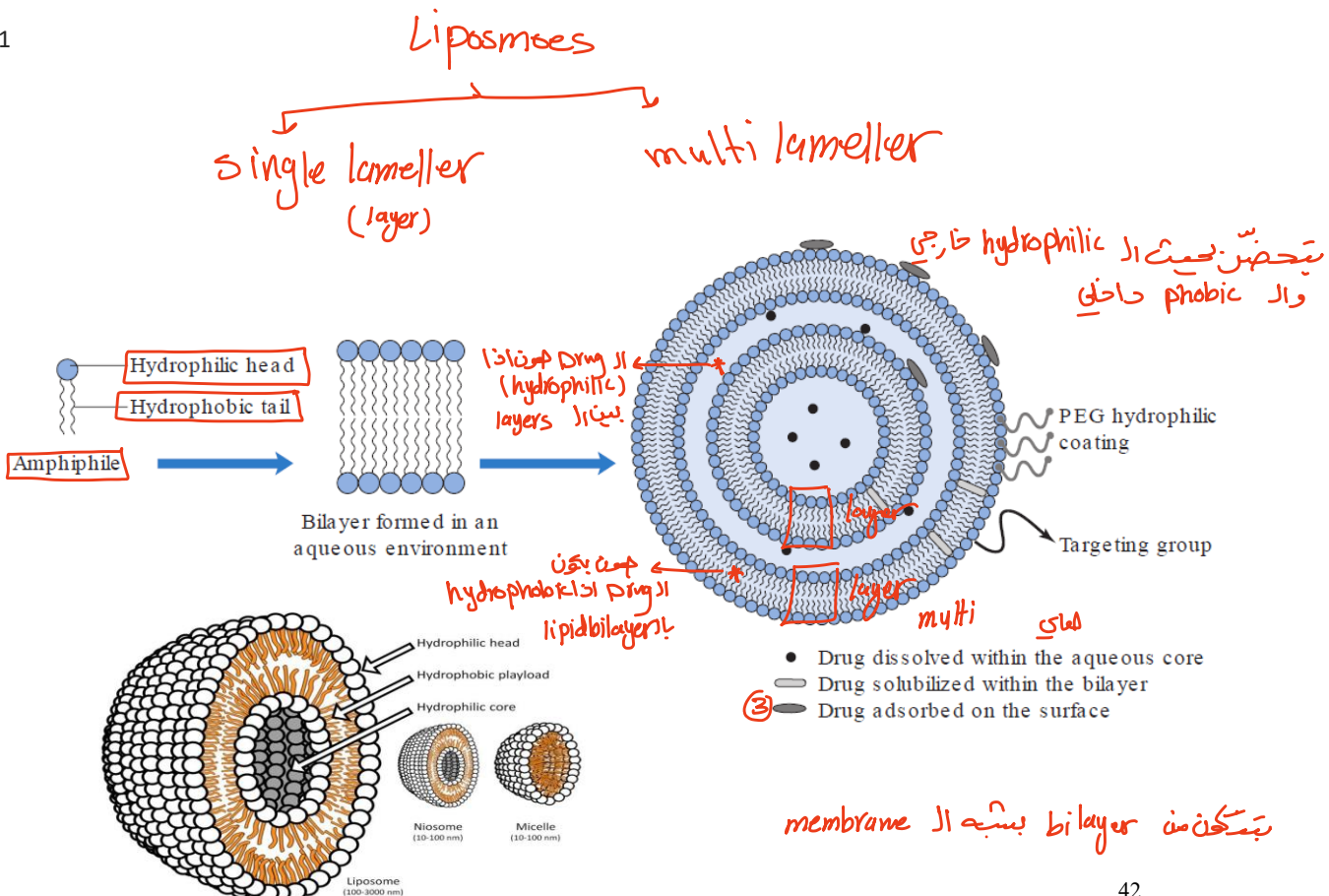
Liposomes and bilayer vesicles

- Liposomes form when the lipids (which are surface active with a hydrophilic head group and a hydrophobic chain(s) at opposing ends) are exposed to an aqueous environment.
 يتم تشكيل micelles
- Under appropriate lipid-to-water ratio and temperature, the lipids will arrange into bilayer vesicles.
 ad T حبراً صفة لا stability
- Unlike micelle formulation, which form spontaneously, energy must be added to the system to drive the formation of liposomes.
 ad T حبراً صفة لا stability

عشان اكون liposomes استخدم سرعة عالية / ال energy متوزعة ولما نتجاوز ال CMC يتكون ال micelle .
 micelle spontaneously , في البداية يتكون ال micelle
 critical micelle concent

41

41



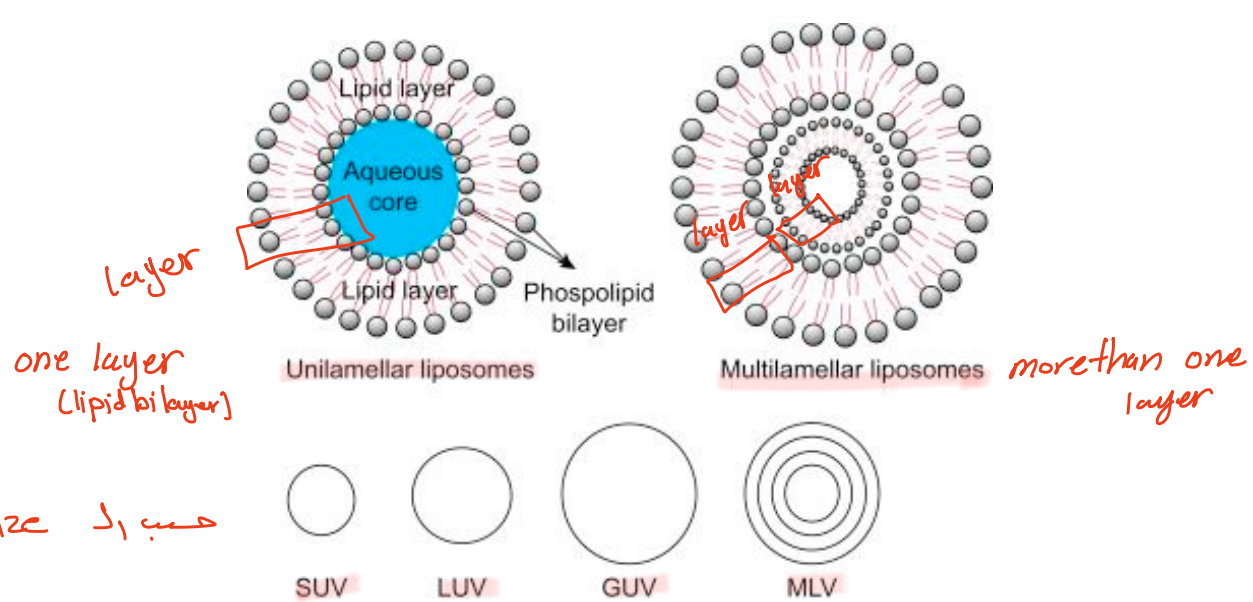
42

42

* يعني يمكن استخدام liposomes لـ hydrophilic + hydrophobic

* بتقطة في لين حكيها Adsorbed من Covalent؟ Covalent بي condition معينة قوية من solvent وحرارة

protection
 Physical Adsorption
 sensitive
 lipid
 * بعض نېچې PEG غاڻ تړيد او half life وځان سټريڪ
 Immunesystem ڏي



S: Single, L: Large G: Giant U: Uni M: Multi, L: Lamellar V: Vesicle
 43

43

Liposomes and bilayer vesicles

- Due to their nature, liposomes are able to carry both water-soluble and lipophilic moieties, with the water-soluble drugs being incorporated within the aqueous compartments and lipid-soluble drugs incorporated within the bilayer.
 - aqueous part ڏي
 - lipid ڏي bilayer
 - Adsorption ڏي drug
- In addition, some drugs and molecules can be adsorbed onto the surface of the liposomes through electrostatic interactions.
- Liposomes can be formulated in a range of diameters from around 30 nm up to several micrometres, therefore they can be considered as nanotechnology.

44

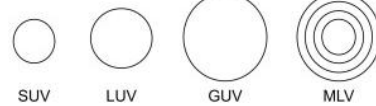
44

Liposomes and bilayer vesicles

Small unilamellar vesicles (SUV)

- These are single-bilayer vesicles, around 30 to 100 nm in size.
- These are generally easier to prepare in a homogeneous size range compared to other types of vesicles and are the most commonly used in clinically approved products.
- Due to their small size there is a low ratio of internal aqueous volume per mole of lipid.

لما انه جعما صغير فال Aqueous core صغير



Addition خلال التحضير نتحكم بار ratios = طريقة الـ size اكبر بتعني

45

45

Liposomes and bilayer vesicles

Large unilamellar vesicles (LUV)

- These are large single-bilayer vesicles of 100 nm and greater.
- These vesicles offer a larger aqueous compartment compared with SUVs.

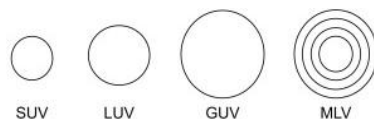
Aqueous core اكبر من الجوز قبل

Multilamellar vesicles (MLV)

- These vesicles have multiple concentric bilayers and are 100 nm to several micrometers in size, depending on their composition and their method of preparation.

حسب طريقة التحضير

- Their low aqueous volume (due to the multiple bilayers) reduces their capacity or carrying water-soluble drugs. [



لانه الداخل منه lipid

فال Aqueous قليل فال

الـ بتقدر تحمله اقل

w. soluble drug

46

46

موجود بالسوق

Liposomes and bilayer vesicles

Examples on clinical applications

الاسماء التجارية من محلولية



1. Cancer chemotherapy (e.g. Caelyx[®] and DaunoXome[®])
2. Treatment of systemic fungal infections (e.g. AmBisome[®])
3. Delivery of vaccines (e.g. Inflaxal V[®])
4. Sustained drug release (DepoCyte[®] and DepoDur[®])
morfine



اي محمد قرأته ما يعرف اذا كانه حفظ

الاسماء التجارية X

Examples of clinically approved liposome products

Product name	Drug	Formulation	Clinical indications
AmBisome [®]	Amphotericin B	SUV liposomes. The vesicles are less than 100 nm in size. Amphotericin B is intercalated within the liposome membrane.	An antifungal agent given intravenously for the treatment of systemic fungal infections
DepoCyte [®]	Cytarabine <i>chemo</i>	Multivesicular vesicles. The vesicles are 3 to 30 μm in size. Cytarabine is entrapped in the aqueous compartments of the liposomes. <i>small</i>	Intrathecal treatment of lymphomatous meningitis.
Myocet [®]	Doxorubicin <i>small + single layer</i>	SUV liposomes around 45 nm in size. Doxorubicin is loaded into the aqueous core of the liposomes where it forms a doxorubicin citrate complex	First-line treatment of metastatic breast cancer in women.

micelle ال liposomes ال

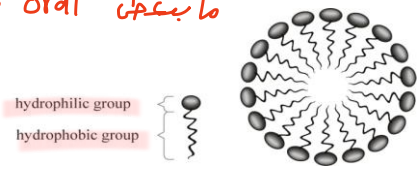
Micelle systems

- Micelles are widely used to formulate low solubility drugs in colloidal solutions.
- Their size (often < 100 nm) means that micelles can be considered within the nanotechnology classification.
- Fugizone® is a mixed micellar formulation which is employed to solubilize Amphotericin B, an antifungal agent used to treat invasive fungal infections, such as systemic candidiasis and histoplasmosis.

very very hydrophobic Drug

hydrophobic drug جوار hydrophobic core tail

parenteral oral ما ينطبق



Amphiphilic part ال liposomes ال micelles ال

Surfactant تتحضر من lipids ال micelles تتحضر من

Dendrimers

راحتة على 52 بعدين رجعت جون

- ❖ Dendrimers are highly branched polymeric, star-shaped macromolecules which can be prepared in the nanosize range.
- ❖ There are three main elements to dendrimers:
 1. A central core نقطة البرارة
 2. The internal dendritic structure, which is composed of the branched polymeric structure built onto the central core
 3. The exterior surface of the dendrimer.

نتم تحضيرها ب nanosize

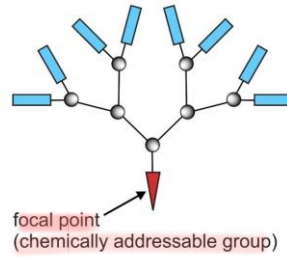
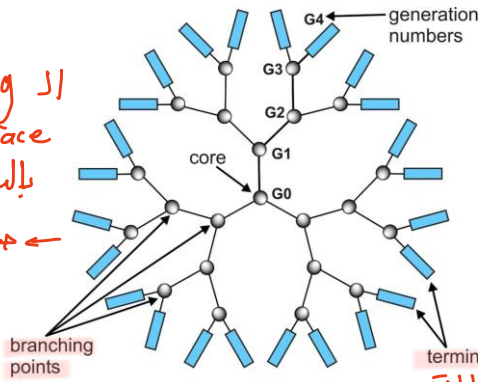
بترتيبهم وحدة وحدة زو Generations

معنى ال conjugation Drug ال عليه ال PEG

These branched polymeric structures are synthesized by step-wise addition of layers of polymer branching, referred to as generations (termed G1, G2, etc.)

زي تفويات الشجرة ، بتحضّر by reactor يحن مفاعل chemical

ال Drug مكن يكون عال surface و مكن بين ال branches بالداخل
 ← مكن ال Drug عال surface



4 Generations هي DENDRIMER
 G0 G1 G2 G3 G4 generations يسورها

DENDRON basic unit

ممكن اعلم ال Dendrimer exterior

A dendrimer (coloured) with 4 generations labelled G1 to 4 respectively. Drug molecules and targeting groups can be conjugated to the exterior of the dendrimer. PEG can also be added to the exterior surface to provide a hydrophilic coating.

51 دمكن تزيد ال half life و تزيد ال stability

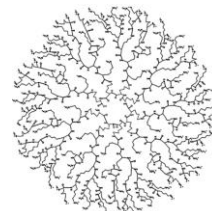
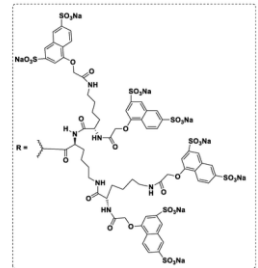
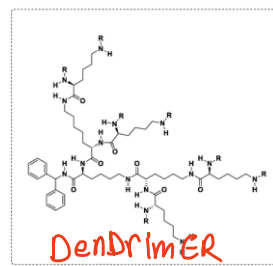
ال Focal point بئال G0 (generation zero) حد ال core لحد بنبلش نهيته

علما chemical Reactions by باقي ال Generations

- مكن اهنى PEG

Dendrimers

- **VivaGel®** is a microbicide gel which uses dendrimers.
- In this formulation, the dendrimer is the **active ingredient** in its own right
- The dendrimer has antiviral properties due to its ability to **bind to viruses** and thereby **blocking their ability to infect cells**.



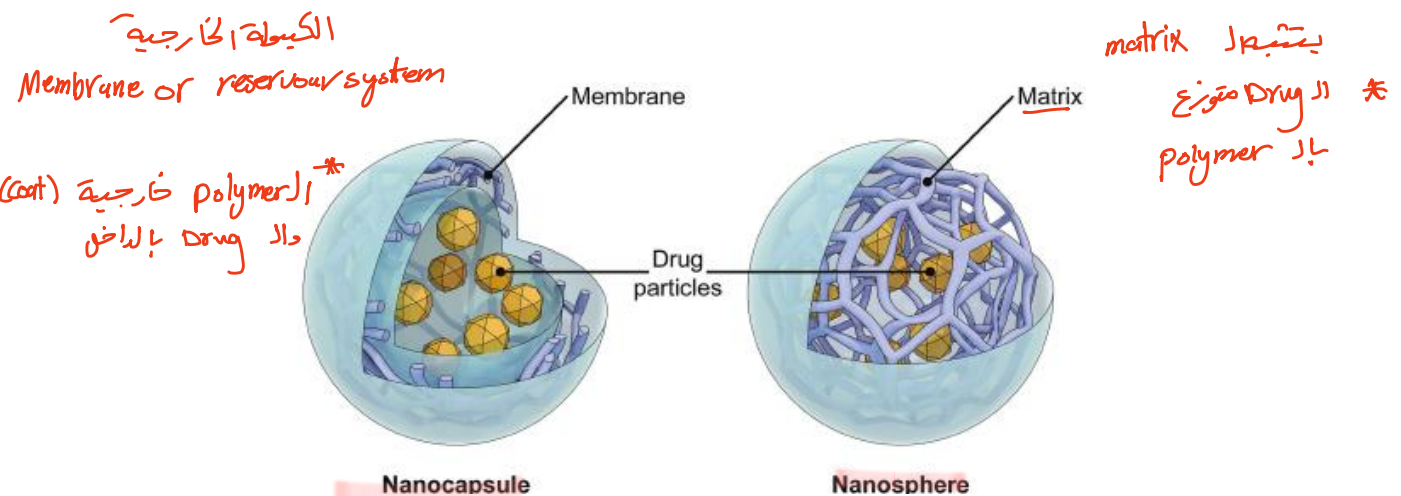
ال generation

Nanoencapsulation

- Nanoencapsulation of drugs involves forming drug loaded particles with diameters ranging from 1 to 1000 nm.
- Owing to their small size, NPs exhibit interesting properties, making them suitable for a variety of drug delivery applications.
- One can distinguish two types of NPs:
 - nanomatrices, which are matrix systems (some times termed also nanospheres); and *monolayeric*
 - nanocapsules, which are reservoir systems composed of a polymer membrane surrounding an oily or aqueous core.

53

53



Schematic representation of a nanocapsule (left). nanomatrix (right)

In Nanocapsules, a core-shell structure with a liquid core surrounded by a polymer shell.
In nanospheres (matrix), the whole particle consists of a continuous polymer network.

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54

بمقتضىها بس ال Nano sphere اخرج بكثر (تأخذوا)