

PETROPHYSICS

Physical properties of natural stone
and other porous mineral materials

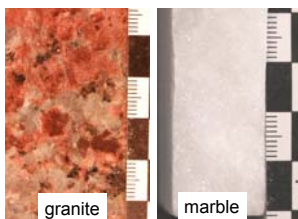
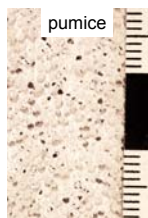
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Depending on the use
(cobblestone, stair, façade,
carved element, pile of bridge...),
stone must have different
physical characteristics.

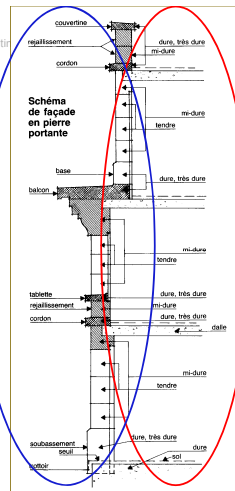
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The physical properties
of stones depend on
cementation, porosity...



they depend on the
origin of stones

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Très dure: very hard
Dure: hard
mi-dure: half-hard
tendre: soft

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[illegible]

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Petra monastery, Jordan

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Interactions between stones and conservation products are also strongly dependent on quantity and quality of the porous space



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Conservation products for stones*, mechanical and/or chemical cleanings -as well as water and salts- can change the physical properties of stones

* (consolidants, water-repellents, anti-graffiti products, paints, ...)

(Total) porosity

Definition: fraction of the total volume of a material « occupied » by voids

$$Pt (\%) = 100 \times \frac{V_v}{V_t} = 100 \times \frac{V_v}{V_v + V_s}$$

Pt = total porosity

Vv = volume of the voids

Vt = bulk volume

Vs = volume of solid

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Densities and porosity

Density (*masse volumique*): quantity of solid matter contained in a given volume of material

$$\text{bulk density } \rho_a = \frac{m}{V_t}$$

(m. vol. apparente)

$$\text{particle or true density } \rho_s = \frac{m}{V_s}$$

(m. vol. réelle ou solide)

$$Pt (\%) = 100 \times \left(1 - \frac{\rho_a}{\rho_s} \right)$$

If Pt tends to 0, ρ_a tends to ρ_s

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Different kinds of porosity

- **Primary** (*primaire*): structural arrangement existing since the genesis of the stone
- **Secondary** (*secondaire*): results from the phenomena of deterioration, diagenesis, metamorphism ...
- **Connected**: voids communicating freely between them
- **Occluded**: not inter-connected voids
- **Intergranular**: voids between the constituent grains
- **Intragranular**: voids inside the constituent grains
- **Micro- and macro-porosity**: < or > to 7.5 μm (from mercury injection porosimetry)

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Different kinds of porosity

Porosity and water retention in soils

Pores	diameter	Water
Big pores	> 50 μm	free water - dry quickly and easily (no capillary action)
Medium pores	10 to 50 μm	free/capillary water - take time to the dry out ("coarser" part of the capillary porosity)
Fine pores	0.2 to 10 μm	capillary water - slow to dry, high capillary tension
Very fine pores	< 0.2 μm	linked (adsorbed) water - very difficult/impossible to dry

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Total porosity ranges for various natural rocks

Rock type	Porosity (%)
sandstones	5 – 50
limestones	5 - 55
crystalline rocks	0 – (10)
volcanic rocks	0 – (90)
metamorphic rocks	0 – (50)

Sources: Freeze and Cherry (1979); McWorter and Sunada (1977).

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Total porosity ranges for various natural swiss rocks

PIERRES NATURELLES	Masse volumique apparente t/m ³	Résistance à la compression N/mm ²	Résistance à la flexion N/mm ²	Porosité totale (connectée) Vol. %	Porosité capillaire Vol. %	Dilatation hydrique mm/m
R. MAGMATIQUES						
Plutoniques	2.6-3.0	80-300	10-40	0.4-1.5	0.2-1.2	0.06-0.2
Volcaniques effusives	1.6-3.1	10-400	5-60	0.2-30	0.1-25	0.06-0.4
R. SEDIMENTAIRES						
Calcaire/ dolomite dures	2.6-2.8	50-200	3-30	0.6-2	0.5-1.5	0.09-0.16
Travertin / tuf calcaire	1.7-2.5	10-80	3-20	2-25	1.5-20	
Calcaire tendre	1.5-2	8-25	2-8	5-25	5-20	0.3-0.5
Grès à ciment siliceux	2-2.6	40-250	7-33	1-25	1-20	0.3-0.6
Grès à ciment calcaire	2-2.6	30-180	3-18	2-20	1-18	0.4-0.8
Conglomérats	2.4-2.6	14-160	2-12	0.5-5	0.5-4	0.1-0.2
R. METAMORPHIQUES						
Gneiss / migmatite	2.5-2.8	70-200	8-45	0.4-2.0	0.25-1.5	
Quartzite	2.6-2.7	100-300	14-60	0.4-2.0	0.2-1.5	
Schistes	2.5-2.7	50-150	8-40	0.6-3.0	0.5-2.5	0.2-0.6
Serpentine	2.6-2.75	55-200	11-60	0.3-2.0	0.25-0.18	
Marbre	2.5-2.7	40-230	10-40	0.3-0.8	0.2-0.5	

D'après Kündig R. et al., 1997. Die mineralischen Rohstoffe der Schweiz. Commission Géotechnique Suisse, Zürich, ETHZ

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(connected) Porosity measurement

The « triple weighing* » method:

M_1 = mass of the dried sample weighing in the air
 M_2 = m. of the water saturated s.**, weighing in the air
 M_3 = m. of the water saturated s.**, weighing underwater

$$M_1 = (\rho_s \times V_s)$$

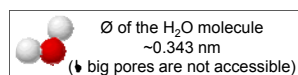
$$M_2 = (\rho_s \times V_s) + (\rho_{H_2O} \times V_v)$$

$$M_3 = (\rho_s \times V_s) - (\rho_{H_2O} \times V_s)$$

avec $\rho_{H_2O} = 1 \text{ g/cm}^3$

$$Pt = \frac{M_2 - M_1}{M_2 - M_3}$$

👉 water soluble phases !!!



*méthode de la triple pesée

** the connected porosity of samples are saturated with water under vacuum

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**(connected) Porosity measurement****The gas pycnometry method:**

Measurement of the volume of a sample (solid or powder). Principle: injection of a gas from a closed volume of reference with a known pressure and a known volume to a closed volume containing the sample. According to the Boyle-Mariotte law, the difference in change of pressure gives the volume of the sample.

V_1 = closed volume of reference

V_0 = volume of the container with sample

V_s = "solid" volume of the sample

P_1 = gas pressure in the reference container

P_2 = gas pressure in the container with sample after expansion of the gas

$$V_s = V_0 - V_1 \left(\frac{P_1}{P_2} - 1 \right)$$

● Ø of the He atom
~0.12 nm ?

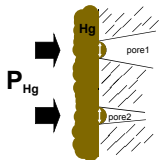
**(connected) Porosity measurement****Mercury intrusion porosimetry (MIP):**

The technique involves the intrusion of mercury, a non-wetting liquid, at high pressure into a material through the use of a porosimeter. The pore size can be determined based on the external pressure needed to force the liquid into a pore against the opposing force of the liquid's surface tension (at 25° C $\gamma = 0,474$ N/m and the contact angle $\alpha = 130^\circ$)

$$P_{Hg} - P_{vap} = \frac{2\gamma \cos\alpha}{R} \quad \text{as } P_{vap} \ll P_{Hg} \Rightarrow P_{Hg} = \frac{2\gamma \cos\alpha}{R}$$

👉 deformable materials ?

● Ø of the Hg atom 0.3 nm
BUT the Ø of the smallest accessible pores depends on the max. P_{Hg} (and the largest ones depends on the min. P_{Hg})

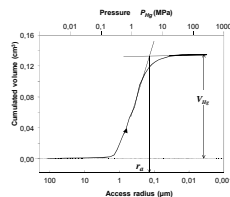
**(connected) Porosity measurement****MIP:**

With a known P_{Hg} on a known volume of mercury, all the pores with an access radius $\geq R$ can be filled

As $P_{Hg} \nearrow$, the radius R of the pores that can be filled \searrow

V_{Hg} = bulk volume of intruded mercury ($\approx Pt$)

ra = threshold radius (the smallest radius giving access to the maximum porous volume)





Speed of sound, porosity and cementation

The time of transmission of the P waves (longitudinal) through the thickness of a porous material \nearrow when:

- the percentage of the void volume \nearrow
- the cementation degree \searrow

The Hooke's law (very) simplified allows to estimate the modulus of elasticity or Young's modulus (E):

$$V_p = \sqrt{\frac{E}{\rho_a}} \Rightarrow \text{relative value}$$



Microscopic observation of thin sections of rocks (polarizing petrographic microscopy)

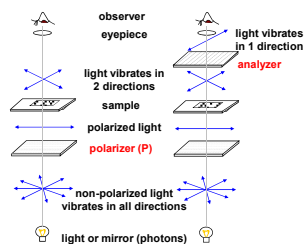
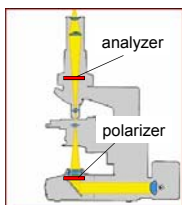
A thin section (*lame mince*) is a 30 μm thick strip of stone which is stuck on a glass slide. This sample can be observed with transmitted light because of its transparency.

Stages of manufacture:

- sawing a stone to « sugar cube » (3 x 2 cm) size
- hardening using an epoxy resin
- polishing of one side which is then glued on the glass slide
- sawing, grinding, then polishing up to a thickness of 30 μm

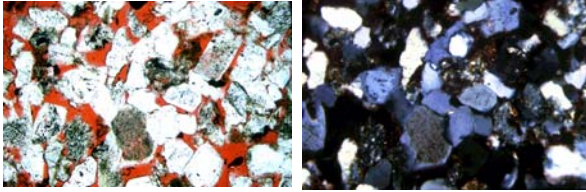


The polarizing petrographic microscope



The optical properties of the minerals in a thin section alter the colour and intensity of the observed light. Minerals can be identified but porosity can also be observed

Microscopic observation of thin sections of rocks



(Photos: D. Jeannette) Sandstone of the Basel cathédrale. Width = 5 mm

- ⇒ mineralogy
- ⇒ geometry of the porous network (only for voids $>1 \mu\text{m}$)

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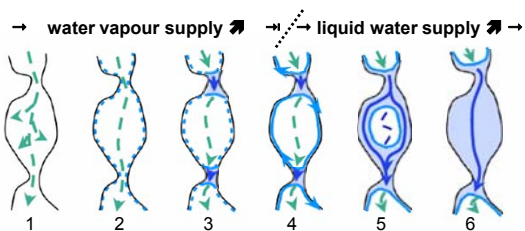
Notion of climate:

In french
http://www.c2rmf.fr/documents/Climat_fiche1.pdf

In English :
<http://www.conservationsphysics.org/atmcalc/atmcalc1.php#svc>

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Water transfer in porous media

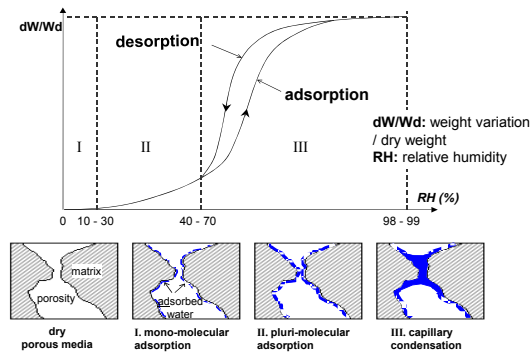


- 1 - vapour diffusion; 2 - vapour adsorption + diffusion; 3 - capillary condensation; 4 - capillary migration along the walls (beginning of filling); 5 - hydraulic flow in unsaturated media (filling in progression or finished); 6 - hydraulic flow in saturated media

According to Rose (1963)

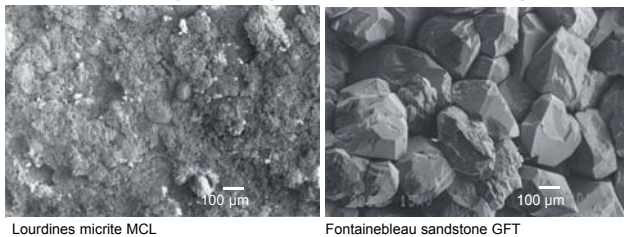
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Water vapour in porous media: adsorption



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Water vapour in porous media: adsorption



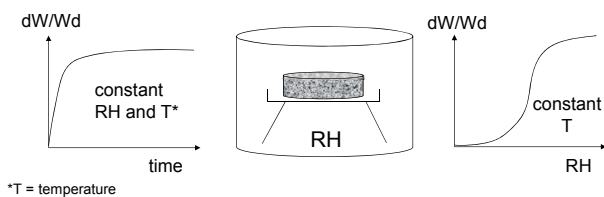
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Hygroscopicity increases when:

- specific surface \nearrow (microporosity, content of clays – for example MCL hygroscopicity > GFT one)
- content of soluble salts \nearrow

Water vapour in porous media: adsorption

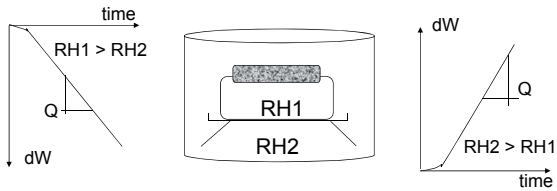
To determine the kinetic adsorption and the adsorption isotherm, samples are first dried at 60°C then, they are placed in closed boxes where relative humidity is controlled. Then, they are weighted regularly until their mass becomes constant.



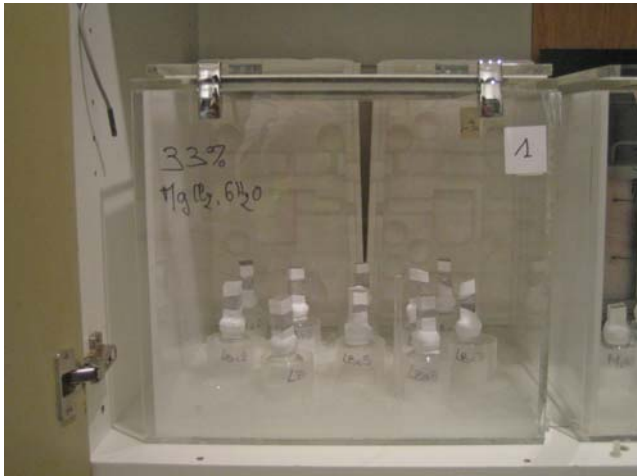
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Water vapour in porous media: permeability

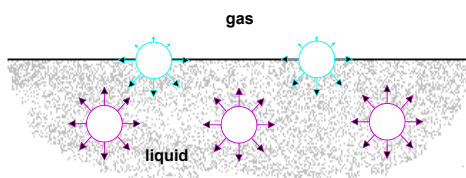
To determine the water vapour diffusivity or the permeability to water vapour samples are first dried at 60°C. Then they are packed in such a way that two opposite faces can exchange with two different relative humidities.



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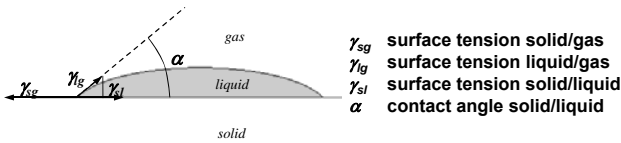
Liquid water in porous media – surface tension



The cause of surface tension in a liquid: a molecule in contact with a neighbour is in a lower state of energy than if it weren't in contact with a neighbour. The boundary molecules have fewer neighbours than interior molecules and are therefore in a higher state of energy than interior molecules => to minimize its energy state, a liquid must minimize its number of boundary molecules and therefore minimize its surface area => formation of "film" => formation of a drop ...

Materials and conservation of built cultural heritage – PhotoPhysics – BR 230

Liquid water in porous media – surface tension



When a drop of liquid is deposited on a solid surface, the interface geometry depends on the affinity of each phase (solid, liquid, gas) to the other, or on their relative surface tensions

Young-Dupré equation:

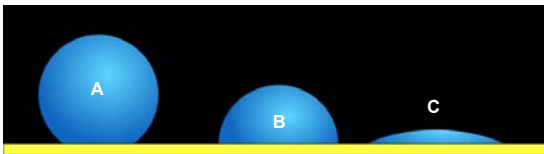
$$\gamma_{lg} \cos \alpha = \gamma_{sg} - \gamma_{sl}$$

wetting liquid: $\alpha < 90^\circ$

non-wetting liquid: $\alpha > 90^\circ$

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Liquid water in porous media – surface tension



Wetting of different fluids. **A** shows a fluid with very **high surface tension** (and thus little wetting), while **C** shows a fluid with **very low surface tension** (more wetting action.) A has a high contact angle, and C has a small contact angle.

Examples: drops of oil, water and water + detergent

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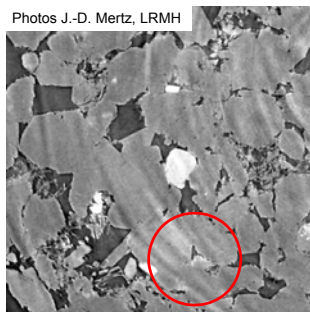
Liquid water in porous media – surface tension

For example, inorganic salts increase the surface tension of a solution, but alcohols or surfactants decrease the surface tension of the solutions

Some treatment effects:

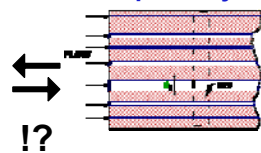
Hydrophobic and consolidant treatments decrease the wettability of the façade surfaces whereas cleaning products tend to increase this wettability

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Buntsandstein sandstone
« Grès à Meules »
Micro computed tomography (x ray - synchrotron)
slice thickness ~1,35 µm

Modeling of the rock porosity



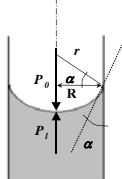
Despite the complexity of the porous structures, it seems that a bundle of parallel capillaries, all of different diameters, embedded in a solid, is a perfect morphology model of a porous medium !!!

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Liquid water in porous media – capillarity

If a tube is sufficiently narrow and the liquid adhesion to its walls is sufficiently strong, surface tension can draw liquid up the tube in a phenomenon known as capillarity.

In such a case, surface tensions are responsible for the capillary pressure (a tension in fact) and the formation of a meniscus at the liquid interface



$$\Rightarrow P_c = P_0 - P_1$$

Laplace equation:

$$P_c = \frac{2\gamma}{r}$$

- P_c = capillary pressure (= tension !)
- P_0 = pressure in the gas
- P_1 = pressure in the liquid
- r = radius of curvature of the meniscus
- R = radius of the capillary
- γ = surface tension liquid/gas

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Liquid water in porous media – capillarity

When the radius of the capillary R is very small, the meniscus is hemispherical and the radius of curvature r can then be connected to R :

$$r = \frac{R}{\cos\alpha} \Rightarrow P_c = \frac{2\gamma \cos\alpha}{R}$$

The more wetting fluid moves spontaneously the less wetting fluid towards the outside of the capillary: this is called a capillary imbibition, suction or absorption

(When a pressure is applied on the less wetting fluid, the more wetting fluid is forced out: this is called a drainage; the MIP for ex.)

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Liquid water in porous media – capillarity

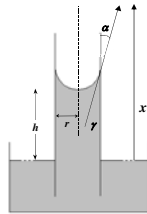
When a capillary tube is plunged in a liquid, the meniscus moves along the tube.

The kinetic follows the **Poiseuille law** which is (if acceleration due to gravity is neglected) :

$$Q(x) = \frac{\pi \gamma \cos \alpha r^3}{4 \eta x}$$

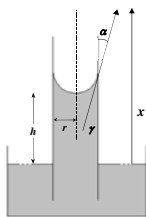
=> when $r \nearrow$, $Q \nearrow$

γ = surface tension liquid/gas
 r = radius of the capillary
 η = dynamic viscosity
 α = contact angle solid/liquid



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Liquid water in porous media – capillarity



When a capillary tube is plunged in a liquid, the meniscus moves along the tube.

The height h of the liquid column at steady state is given by the Jurin's equation :

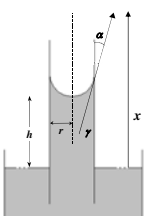
$$h = \frac{2 \gamma \cos \alpha}{r g \rho}$$

=> when $r \nearrow$, $h \searrow$

γ = surface tension liquid/gas
 g = gravity
 ρ = density of liquid
 r = radius of the capillary
 α = contact angle solid/liquid

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Calculation examples



For a water-filled glass tube in air at sea level,

$$\gamma = 0.0728 \text{ J/m}^2 \text{ at } 20^\circ\text{C}$$

$$\theta = 20^\circ (0.35 \text{ rad})$$

$$\rho = 1000 \text{ kg/m}^3$$

$$g = 9.8 \text{ m/s}^2$$

=> the height of the water column is given by:

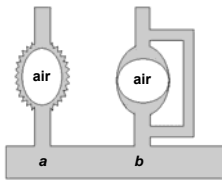
$$h = \frac{2 \gamma \cos \alpha}{r g \rho} \approx \frac{1.4 \times 10^{-5}}{r}$$

Thus for a 2 m wide ($r = 1 \text{ m}$) tube, the water would rise an unnoticeable 0.014 mm.

For a 2 cm wide tube ($r = 1 \text{ cm}$), the water would rise 1.4 mm and for a capillary tube with $r = 0.1 \text{ mm}$, the water would rise 14 cm.

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Liquid water in porous media – capillarity

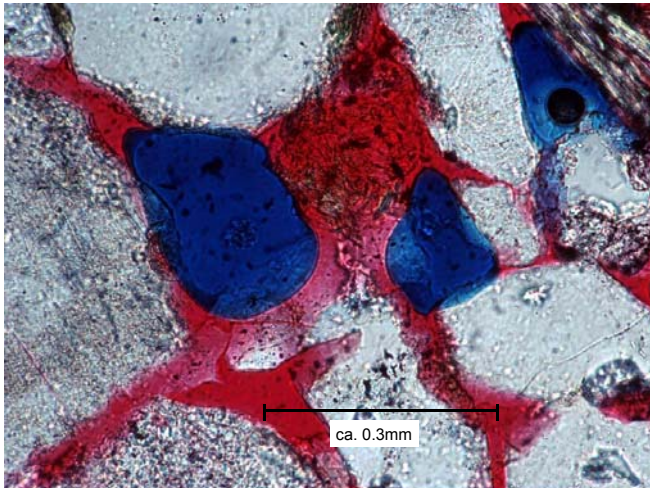


Less wetting fluid trapped by the more wetting fluid during an imbibition because of:
a. rugosity
b. derivation

The porosity of a rock (or a mortar, a concrete,...) is a complex system formed by inter-connected wide and tiny pores. As a result, there are always many parallel ways for the capillary water during an **imbibition**.

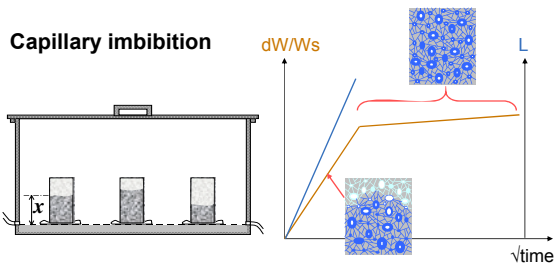
This great complexity allows the water (more wetting fluid) to trap air (less wetting fluid) in parts of the porosity.

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Liquid water in porous media – capillarity

Capillary imbibition

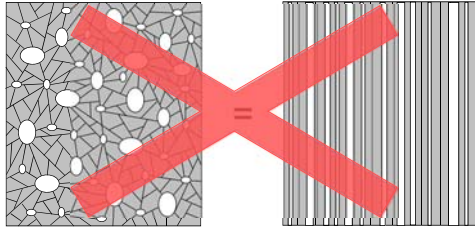


The kinetics of linear and mass capillary imbibitions of a stone show the complexity of the porous network

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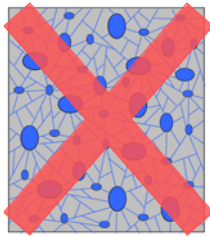
Liquid water in porous media – capillarity



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Liquid water in porous media – capillarity



In a porous network partly formed of (connected) macropores, the water saturation of the porosity is not reached by simple capillarity

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Liquid water in porous media – capillarity

During a **capillary imbibition**:

Porosity freely accessible to water = **free porosity**
 Portion of porous space inaccessible = **trapped porosity**

The trapped porosity is characterised by the capillary saturation coefficient or **Hirschwald coefficient (S%)**

$$S\% = \frac{P_i}{P_t} \times 100 \quad \text{where} \quad P_i (\%) = \frac{M_i - M_1}{M_2 - M_1} \times 100$$

P_i = porosity filled by water imbibition
 M_i = sample weight after water imbibition
 M_1, M_2, M_3 : see page 15

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Liquid water in porous media – gelivity

The ability to be more or less filled with water in the presence of air **is one of the lot of parameters** controlling the frost resistance of a stone

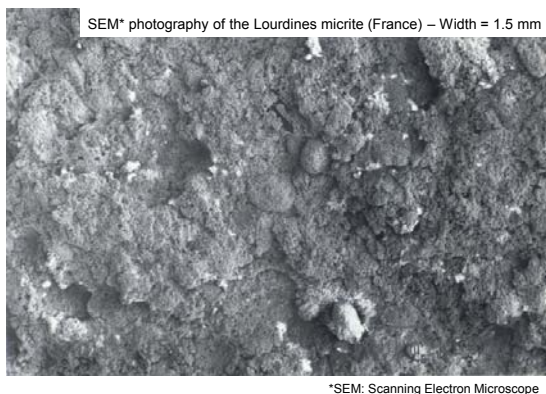
Estimation of frost resistance :

In a “simplistic” way, a stone is described as

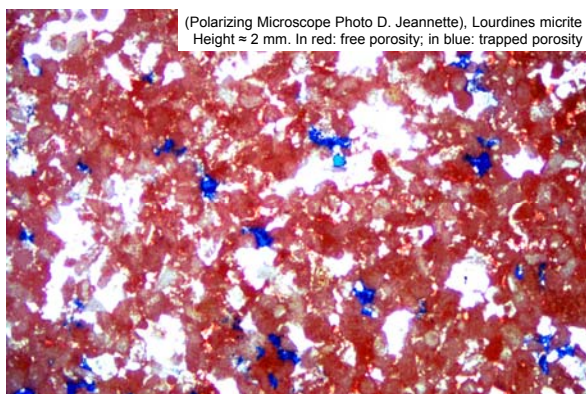
- **frost susceptible** if S% is > 0,85
- **not frost susceptible** if S% is < 0,75

A high water saturation makes the stone frost susceptible (because of the volume expansion of about 9% - or a linear expansion of 3% - of this liquid when it freezes)

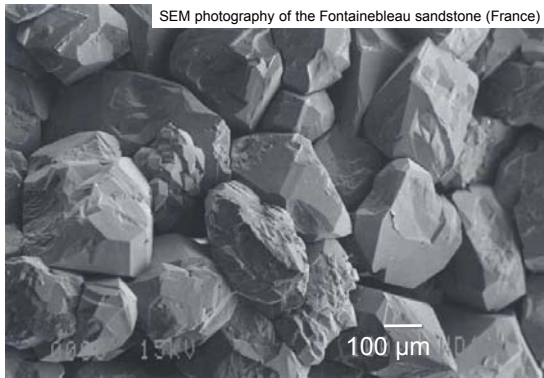
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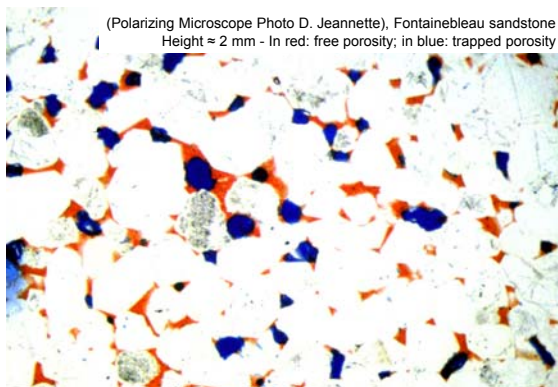
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Materials and conservation of built cultural heritage – Petrophysics – BR 48



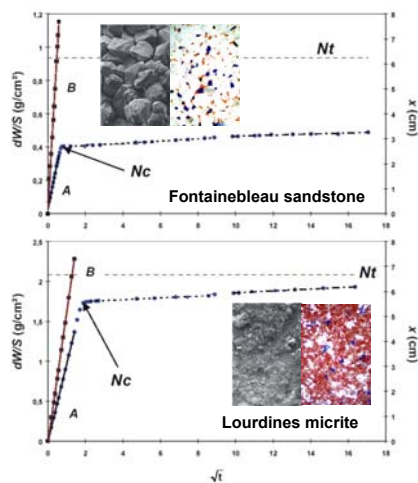
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Materials and conservation of built cultural heritage – Petrophysics – BR 5/0

Capillary imbibition
(RH 100%, T 23° C)

dW = weight variation
S = imbibition surface
Nc = capillary porosity
Nt = total porosity



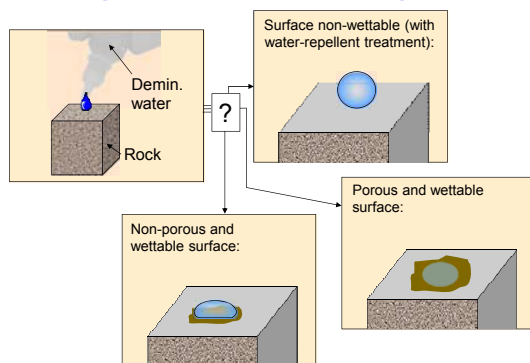
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Liquid water in porous media – capillarity

- In **general**, the higher the dimension of the pores, the higher is the imbibition velocity
- if the linear capillarity is much more rapid than the massic one, the trapped porosity is high => even if water goes up high, the water saturation remains low
- the more varied the pore sizes, the bigger is the trapped porosity: media with very homogeneous porosity can reach high water saturation

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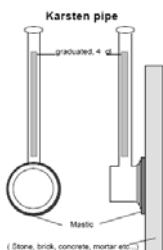
How to quickly estimate the wettability and the water absorption in the field: the drop of water- test



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How to estimate the initial water absorption in the field: Karsten pipes

<http://www.reedconstructiondata.com>



Final Report for the Research and Development Project Non-Destructive Field Tests in Stone Conservation Literature Study Rapport från Riksantikvarieämbetet 2006:3 // <http://www.raa.se/publicerat/9172094345.pdf>

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Conservation Science Consulting Sàrl

How to estimate the initial water or alcohol absorption in the field: Mirowski pipes





Conservation Science Consulting Sàrl

How to estimate the initial water absorption in the field: contact sponge



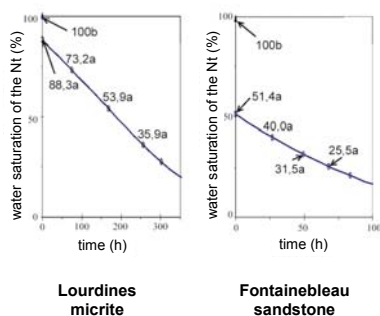
Marini P., Bellopede R., 2009- Bowing of marble slabs: Evolution and correlation with mechanical decay. Construction and Building Materials, Vol. 23, Issue 7, pp 2599–2605

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Liquid water in porous media – drying

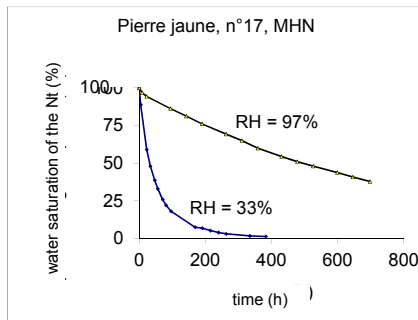


Drying of a specific material is always much slower than its capillary imbibition

In general, the bigger the dimension of the pores of a material, the faster it dries

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Liquid water in porous media – drying



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Materials and conservation of built cultural heritage – Petrophysics – BR 6/9

Liquid water in porous media – dilation (dilatation)

Every porous material changes more or less its dimensions when it adsorbs (vapor) or absorbs (liquid) water (because of a loss of cohesion between grains and or the swelling of clays). This phenomenon is called **hydric dilation** (with liquid water) or **hygric dilation** (with water vapour).

Hydric dilation measurement: the dried sample (length: L_0) is fixed under a dial test indicator (*comparateur*), then it is entirely immersed in water. The extension which is obtained after 72 hours of immersion allows to calculate the hydric dilation coefficient:

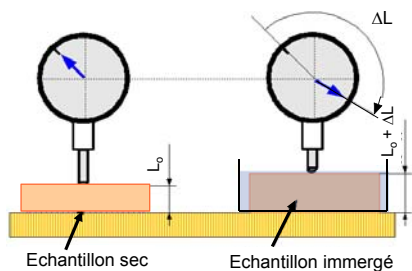
$$\varepsilon(72) = \frac{\Delta L(72 \text{ h})}{L_0}$$



$\varepsilon(72) \geq 2 \text{ mm/m}$ is regarded as high and dangerous

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Liquid water in porous media – dilation (dilatation)



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Notion of R.V.E.

Rocks are **heterogeneous materials** (mineralogy, porosity) on a scale of:

- pores
- samples
- outcrops (*affleurements*)

Therefore to be representative, measurements of physical and mechanical properties of rocks must be done on **Representative Volume Elements** (*volumes élémentaires représentatifs*).

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- <http://www.kasuku.ch/pdf/hygrometrie.pdf>

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