

# **PETROPHYSICS**

Physical properties of natural stone and other porous mineral materials







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

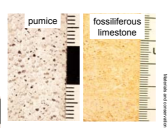


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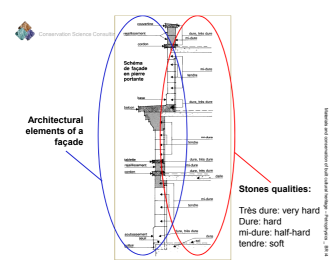
The physical properties of stones depend on cementation, porosity...







◆ they depend on the origin of stones





In the same exposure conditions, the durability of stones lying side by side depends on the liquids and gaseous exchanges between them and their environment (=physical properties)





Exchanges between stones and the environment depend on:
- the quantity of the pore space
- the quality of the pore space (geometry of pores, mineralogical nature of the inner surfaces...)









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







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 x 
$$\frac{Vv}{Vt}$$
 = 100 x  $\frac{Vv}{Vv + Vs}$ 

Pt = total porosity Vt = bulk volume Vv = volume of the voids Vs = volume of solid



## **Densities and porosity**

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

bulk density 
$$\rho_a = \frac{m}{Vt}$$

$$\begin{array}{|c|c|c|c|c|} \hline particle \ or \ true \\ density \\ \text{(m. vol. réelle ou solide)} \end{array} \rho_s = \begin{array}{|c|c|c|c|} \hline m \\ \hline Vs \\ \hline \end{array}$$

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

If Pt tends to 0,  $\rho_a$  tends to  $\rho_s$ 



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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  $\mu$ m (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).



#### Total porosity ranges for various natural swiss rocks

DIEDDES		D/ : /	D/ : /		D "/	D2 1 1
PIERRES	Masse	Résistance	Résistance	Porosité	Porosité	Dilatation
NATURELLES	volumique	à la	à la flexion	totale	capillaire	hydrique
	apparente	compression		(connectée)		
	t/m <sup>3</sup>	N/mm <sup>2</sup>	N/mm <sup>2</sup>	Vol.%	Vol.%	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	
Quarzite	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
Serpentinite	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



#### (connected) Porosity measurement

## The « triple weighing\* » method:

 $\mathbf{M_1}$  = mass of the dried sample weighing in the air

 $\mathbf{M_2}$  = m. of the water saturated s.\*\*, weighing in the air  $\mathbf{M_3}$  = m. of the water saturated s.\*\*, weighing underwater

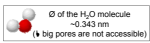
$$M_1 = (\rho_s x Vs)$$

$$M_1 = (\rho_s \times Vs) + (\rho_{H2O} \times Vv)$$
  
 $M_3 = (\rho_s \times Vs) - (\rho_{H2O} \times Vs)$   
 $A_3 = (\rho_s \times Vs) - (\rho_{H2O} \times Vs)$   
 $A_3 = (\rho_s \times Vs) - (\rho_{H2O} \times Vs)$ 

$$M_3 = (\rho_s x Vs) - (\rho_{H2O} x Vs)$$

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

#### ▶ water soluble phases !!!



\*méthode de la triple pesée
\*\*\* the connected porosity of samples are saturated with water under vaccum



# (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<sub>1</sub> = closed volume of reference

 $\mathbf{V_0}$  = volume of the container with sample

 $\mathbf{V_s}$  = "solid" volume of the sample

P<sub>1</sub> = gas pressure in the reference container

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

 $V_s = V_0 - V_1 \left[ \frac{P_1}{P_2} - 1 \right]$ 





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### (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  $25^\circ$  C  $\gamma$  = 0,474 N/m and the contact angle  $\alpha$  =  $130^\circ$ )

$$P_{Hg} - P_{vap} = \frac{2\gamma \cos \alpha}{R}$$

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



Ø of the Hg atom 0.3 nm
BUT the Ø of the smallest
accessible pores depends on the
max. PHg (and the largest ones
depends on the min. PHa



#### (connected) Porosity measurement



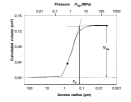
#### MIP:

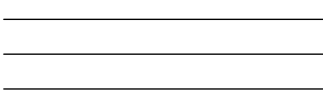
With a known  $P_{Hg}$  on a known volume of mercury, all the pores with an access radius  $\geq R$  can be filled

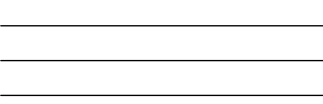
As  $P_{Hg}$  7, the radius R of the pores that can be filled  $\mathbf{3}$ 

 $V_{Hg}$  = 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 7
- the cementation degree \(\mathbb{\sigma}\)

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

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





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# 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 sticked on a glass slide. This sample can be observed with transmitted light because of its transparency.

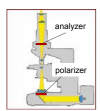
#### Stages of manufacture:

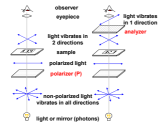
- sawing a stone to « suger cube » (3 x 2 cm) size
- hardening using an epoxy resin
- polishing of one side which is then glued on the glass slide
- sawing, grinding, then polishing up to a thickness of 30 μm

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#### 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

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## Microscopic observation of thin sections of rocks





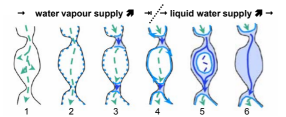
- $\Rightarrow$  mineralogy
- $\Rightarrow$  geometry of the porous network (only for voids >1  $\mu$ m)



Notion of climate: In french http://www.c2rmf.fr/documents/Climat\_fiche1.pdf In English :

# Water transfer in porous media

vapour transfer liquid transfer ..... adsorbed vapour

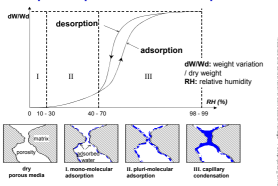


1 - vapour diffusion; 2 - vapour adsorption + diffusion; 3 - capillary condensation; 4 - capillary migration along the walls (beginning of filling); 5 - hydraulic flow in unsaturated media (filling in progression or finished); 6 - hydraulic flow in saturated media

According to Rose (1963)



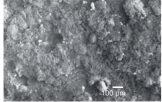
#### Water vapour in porous media: adsorption

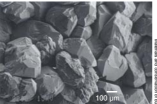




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## Water vapour in porous media: adsorption





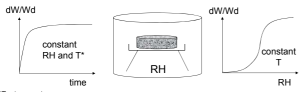
#### Hygroscopicity increases when:

- specific surface **7** (microporosity, content of clays for example MCL hygroscopicity > GFT one)
- content of soluble salts 7



# 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.



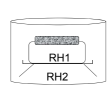
\*T = temperature

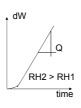


#### Water vapour in porous media: permeability

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.



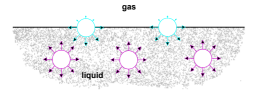








### Liquid water in porous media - surface tension

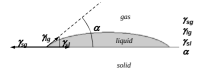


The cause of surface tension in a liquid: a molecule in contact with a neighbour is in a lower state of energy than if it weren't in contact with a neighbour. The boundary molecules have fewer neighbours than interior molecules and are therefore in a higher state of energy than interior molecules => 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 ...

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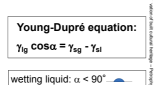


### Liquid water in porous media - surface tension



surface tension solid/gas surface tension liquid/gas surface tension solid/liquid contact angle solid/liquid

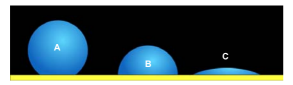
When a drop of liquid is deposited on a solid surface, the interface geometry depends on the affinity of each phase (solid, liquid, gas) to the other, or on their relative surface tensions



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



## 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



#### Liquid water in porous media - surface tension

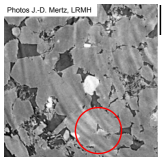
For example, inorganic salts increase the surface tension of a solution, but alcohols or surfactants decrease the surface tension of the solutions

#### Some treatment effects:

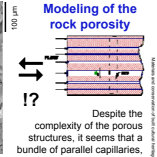
Hydrophobic and consolidant treatments decrease the wettability of the façade surfaces whereas cleaning products tend to increase this wettability

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



all of different diameters, embedded in a solid, is a perfect morphology model of a porous medium !!!



### 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



 $Pc = P_0 - P_1$ 

Laplace equation:

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

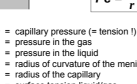
Pc = capillary pressure (= tension !)

= pressure in the gas

= radius of curvature of the meniscus

= surface tension liquid/gas







#### 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 rcan then be connected to R:

$$r = \frac{R}{\cos \alpha}$$



$$Pc = \frac{2\gamma\cos\alpha}{R}$$

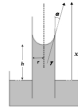
The more wetting fluid moves spontaneously the less wetting fluid towards the outside of the capillary: this is called a capillary imbibition, succion or absorption

(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.)



When a capillary tube is plunged in a liquid, the meniscus moves allong 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 7, Q 7

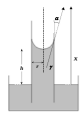
 $\begin{array}{ll} \gamma & = \text{surface tension liquid/gas} \\ r & = \text{radius of the capillary} \end{array}$ 

 $\eta$  = dynamic viscosity

 $\alpha$  = contact angle solid/liquid



# Liquid water in porous media - capillarity



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

The height h of the liquid column at stady state is given by the Jurin's

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

=> when r 7, h ≥

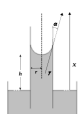
g = gravitiy

= density of liquid

= radius of the capillary  $\alpha$  = contact angle solid/liquid



#### Calculation examples



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

 $\gamma = 0.0728 \text{ J/m}^2 \text{ at } 20^{\circ}\text{C}$   $\theta = 20^{\circ} (0.35 \text{ rad})$ 

 $\rho = 1000 \text{ kg/m}^3$   $g = 9.8 \text{ m/s}^2$ 

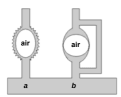
=> the height of the water column is given by:

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

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

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





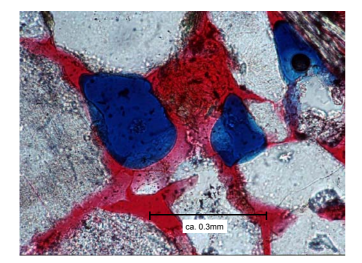
Less wetting fluid trapped by the more wetting fluid during an imbibition because of:

- a. rugosity
- b. derivation

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

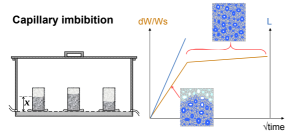
This great complexity allows the water (more wetting fluid) to trap air (less wetting fluid) in parts of the porosity.







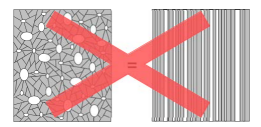
## Liquid water in porous media - capillarity



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

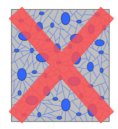
se seden - a conclusioned - passing



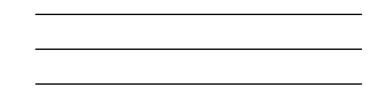




## Liquid water in porous media - capillarity



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





## 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 caracterised by the capillary saturation coefficient or **Hirschwald coefficient (S%)** 

$$S\% = \frac{P_i}{Pt} \times 100$$
 where  $Pi (\%) = \frac{M_i - M_1}{M_2 - M_3} \times 100$ 

 $P_i$  = 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** controling 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

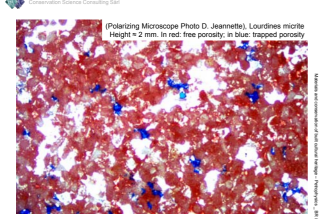
A high water saturation makes the stone frost susceptible (because of the volume expansion of about 9% - or a linear expansion of 3% - of this liquid when it freezes)



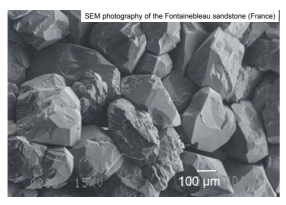


\*SEM: Scanning Electron Microscope

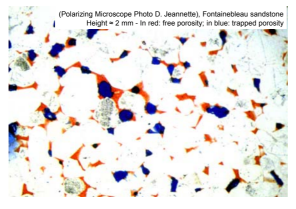


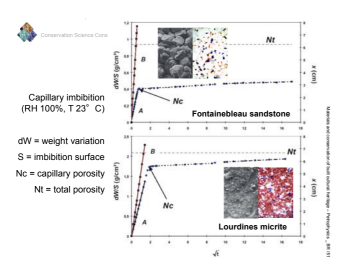












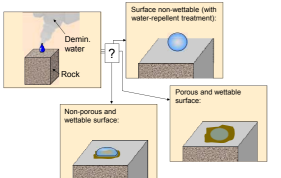


- In general, the higher the dimension of the pores, the higher is the imbibition velocity
- if the linear capillarity is much more rapid than the massic one, the trapped porosity is high => even if water goes up high, the water saturation remains low
- the more varied the pore sizes, the bigger is the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



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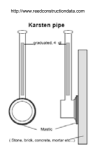
How to quickly estimate the wettability and the water absorption in the field: the drop of water-test



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





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

rials and conservation of built cultural her lage – Petrophysics \_ BR /5



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





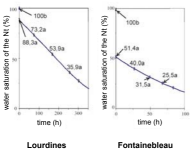
## How to estimate the initial water absorption in the field: contact sponge



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



# Liquid water in porous media - drying

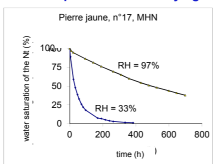


Lourdines micrite sandstone Drying of a specific material is always much slower than its capillary imbibition

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



## Liquid water in porous media - drying







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## 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).

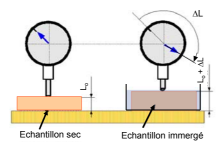
Hydric dilation measurement: the dried sample (length:  $L_0$ ) is fixed under a dial test indicator (comparateur), then it is entirely immersed in water. The extension which is obtained after 72 hours of immersion allows to calculate the hydric dilation coefficient:

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

ε(72) ≥ 2 mm/m is regarded as high and dangerous



## Liquid water in porous media - dilation (dilatation)





#### 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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