Quantitative Real-Time PCR

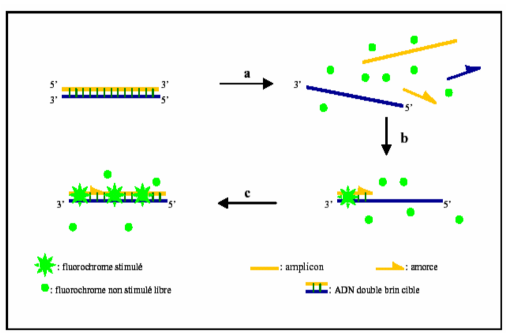
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Real-Time PCR: principle

- 1. The real time PCR is based on the detection and the quantification of a fluorescent transmitter during the process of amplification.
- 2. The increase in the fluorescent signal is directly proportional to the quantity of amplicons produced during the reaction.
- 3. Two general principles for the quantitative detection of amplicons:
 - agents binding to the double-stranded-DNA (SybrGreen I)
 - fluorescent probes (FAM, TAMRA, JOE, ROX,...)
- 4. For the fluorescent probes, there are 4 main technologies:
 - probe hydrolysis
 - hybridisation of 2 probes
 - molecular beacons
 - scorpion primer

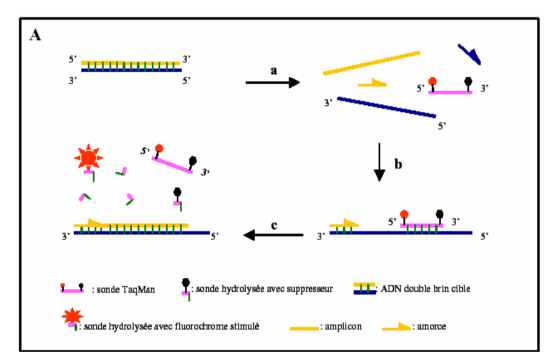
Equivalent sensitivity Different Specificity

Agents binding to the double-stranded-DNA (SYBR Green I)



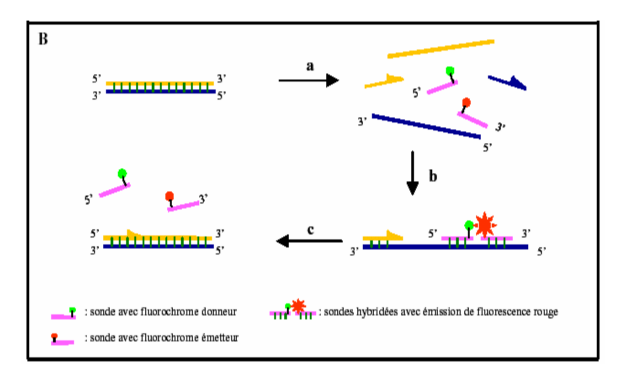
- a. The free SYBR Green exhibits little fluorescence at the time of the denaturation.
- b. With the temperature of pairing, some molecules bind to the nascent double-stranded-DNA.
- c. During the polymerisation step, more and more of molecules bind to the nascent strand and the increase in fluorescence can be followed in real time.

Hydrolysis probes (Taqman)



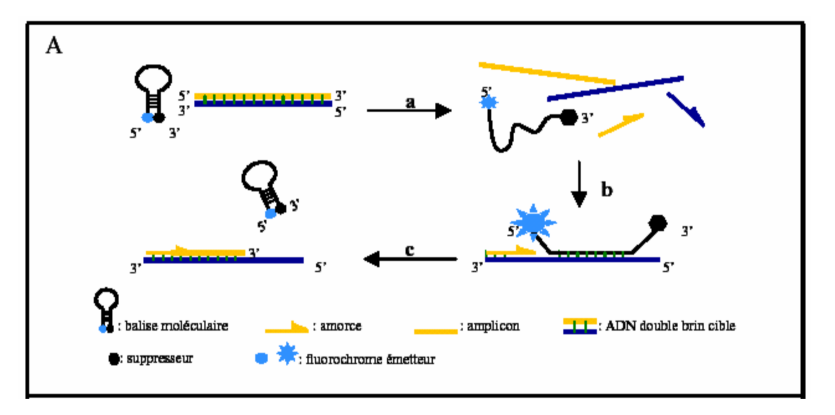
- a. During the denaturing step, the probe is free on solution.
- b. During the annealing step, both probes hybridise to their target sequence. The proximity of the fluorochrome allows the inhibition of fluorescence.
- c. The polymerase moves and hydrolyses the probe. The transmitting fluorochrome is released from the environment of the suppressor thus allowing the emission of fluorescence.

Hybridisation probes (HybProbes)



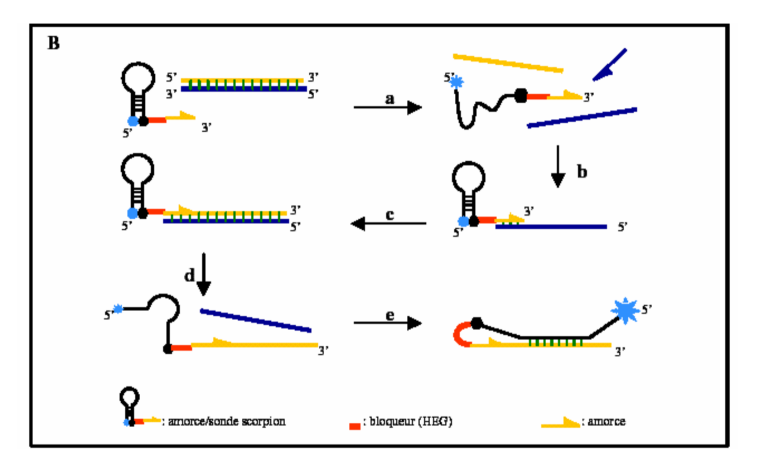
- a. During the denaturing step, in solution the 2 probes are apart.
- b. During the annealing step, both probes hybridise to their target sequence. The proximity of the fluorochrome allows the red emission of fluorescence.
- c. The probes turn over free in solution.

Molecular beacons



- High specificity + high precision
- When T°C<Tm, hairpin structure \rightarrow no hybridisation and no fluorescence

Scorpion primer



- Suppressor HEG prevents the replication of the molecular beacon
- One of the best method

The SYBR Green: advantages and disadvantages

1.<u>Advantages</u>:

- Economic
- Easy to use

- Has more sensibility than the ethidium bromide (another intercalating agent)

- Does not inhibit the reaction of amplification
- Does not require any fluorescent probe, thus does not require any particular expertise for the design of the probes
- Is not affected by mutations in the target DNA

2.Disadvantages:

- Impossible to make sure of specificity of amplicons

- Bad pairing can lead to positive forgeries or an over-estimate of the quantification

- The emission of fluorescence can be skewed by the molecular mass of the DNA amplified by a longer amplicon which will fix more fluorescent molecules compared to a shorter amplicon in the same reaction

- Still unspecified mutagen capacity

Hydrolysis probes: advantages and disadvantages

1. Advantages:

- Increased specificity: the specificity of hybridisation between the fluorescent probe and the sequence of DNA significantly reduces the emission of non-specific fluorescence due to bad pairings or primers dimers.

- Better capacity of multiplexing: reactions multiplex can be elaborate by using distinct transmitting fluorochromes related to different probes in a PCR reaction.

2. Disadvantages:

- Taqman technology is less effective and less flexible device that other technologies in real time for the detection of specific mutations.

- To respect the principles of design of the probes

Probe design

- 1. 30 bases in length
- 2. G-C content of around 50 %
- 3. The probes should not overlap with, or have sequence complementarity with either of the primers
- 4. Probes should not contain a G at their 5' ends, because such an arrangement quenches reporter fluorescence, even after cleavage
- 5. A Tm of 5 to 10°C greater than that of the primers

Hybridisation probes: advantages and disadvantages

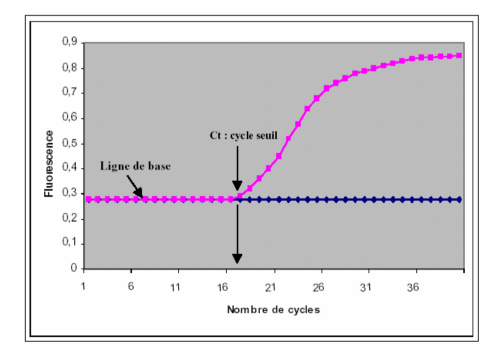
1. Advantages:

- High specificity
- High flexibility for probe design
- As the probes are not hydrolysed, they are used at each cycle
- 2. Disadvantages:
 - Taqman probe design

Threshold cycle = Ct(1)

-The concept of the threshold cycle is at the heart of accurate and reproducible quantification using fluorescence-based PCR.

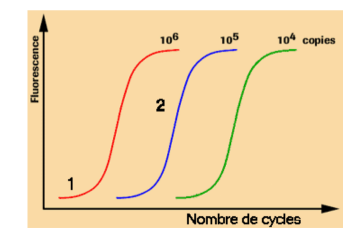
-It corresponds to the cycle from which one observes a statistically significant increase in standardized fluorescence



Threshold cycle = Ct(2)

- The more template present at the beginning of the reaction, the fewer number of cycles it takes to reach a point for which the fluorescence signal is first recorded as statistically significant above background. This point is defined as the Ct.

- The threshold cycle will always occur during the exponential phase of amplification.



Threshold cycle = Ct (3)

-Quantification is not affected by any reaction components becoming limited in the plateau phase

-The Ct value can be translated into a quantitative result by constructing a standard curve

$\mathsf{R} = 2^{-\Delta \mathsf{C} \mathsf{t} 1 / \Delta \mathsf{C} \mathsf{t} 2}$

 Δ **Ct1** = Δ Ct target gene = Ct target gene with treated sample – Ct same gene with sample calibrator

 Δ Ct2 = Δ Ct standardizing gene = Ct standardizing gene with treated samples – Ct same gene with sample calibrator

 \Rightarrow The difference between two samples is considered as significant from one Ct.

Development

