

**NANOCOLOR® TTC/Sludge activity 150****Method:**

Determination of the biochemical activity of sludge (activated sludge, digested sludge etc.) by means of the dehydrogenase activity using 2,3,5-triphenyltetrazoliumchloride (TTC). Colourless TTC is converted into red triphenylformazane (TPF) by dehydrogenases. The formed, water-insoluble TPF is dissolved in ethanol and is photometrically determined.

The rapid test enables:

1. the determination of the biochemical activity ( $A_5$ ) of sludge in terms of the  $\mu\text{g TPF/mg}$  of the dry sludge mass (**method 8901**)
2. the characterization of the effect of waste water and waste water compounds on sludge (stimulation or inhibition of the dehydrogenase activity DHA in percentage) (**method 8902**)
3. a rapid, visual valuation of the degree of stabilisation of sludge based on a simple screening method (**method 8903**)

**Content of reagent set:**

Range:	5 – 150 $\mu\text{g TPF}$ (triphenylformazane) (method 8901)	0.050 – 2.300 E (method 8902)
Factor:	0071.	–
Wavelength (HW = 5 – 12 nm):	470 nm	
Reaction time:	30 – 120 min	
Reaction temperature:	20 – 25 °C	

**Box A:** 20 empty test tubes TTC 150

3 syringes 5 mL

**Box B:** 1 bottle with 30 mL TTC 150 R1

2 bottles with 60 mL TTC 150 R2

2 syringe tubes 5 mL

1 reaction vessel 40 mL

1 screw plug with suction pipe

3 Luer-Lock seal plugs female

2 Luer-Lock connecting adaptors female/female

1 Luer-Lock seal plug male

**Box C:** 21 membran filters  $\varnothing$  1.2  $\mu\text{m}$

**Hazard warning:**

Reagent R2 contains ethanol 90 – 98 %.

For further information ask for a safety data sheet.

**Interferences:**

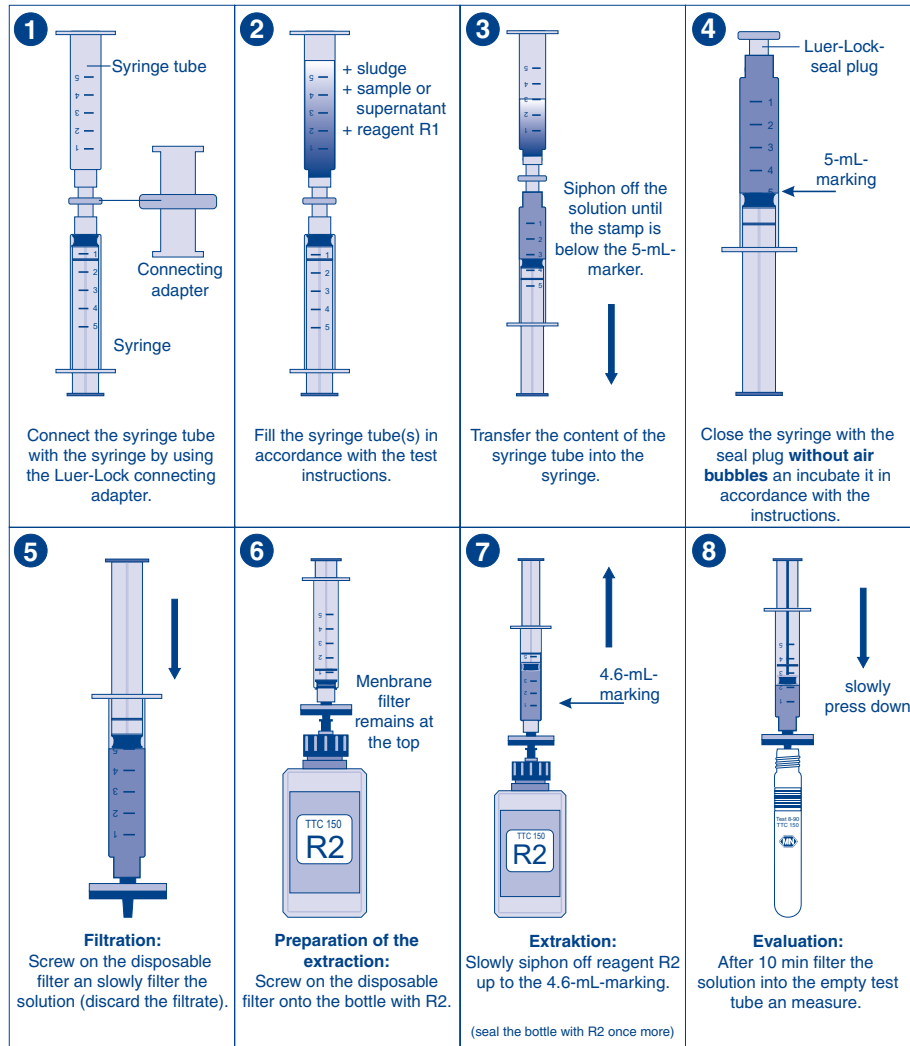
The triphenylformazane (TPF) which is formed is very light sensitive and hence the test samples should be incubated in the dark. Oxygen inhibits the TTC reduction; for this reason the test samples in the syringe tubes should have a constant volume and no oxygen should be enclosed within them. The dehydrogenase activity is not just hindered by oxygen but also by  $\text{NO}_3^-$ ,  $\text{Fe}^{3+}$  and  $\text{NO}_2^-$ . A stimulating effect have  $\text{P}_i$ ,  $\text{Fe}^{2+}$ ,  $\text{SO}_4^{2-}$ ,  $\text{Cl}^-$  and  $\text{Mn(IV)}$ .

**Storage:**

Store the test kit in a cool (2 – 8 °C) and dry place.

**Practical tips:**

1. The result can be influenced by the type and origin of the sludge amongst other things. Hence the origins of the sludge should at least be added to the result. Scum or bulking sludge are not appropriate for routine tests in accordance with method 8902. The processes which have been listed are, however, appropriate for the test of digested sludge.
2. The tests can be carried out at room temperature. Execution at constant, defined incubation temperatures is recommended in order to achieve better comparability of the results of series tests such as hydrograph curves. The incubation of the test samples should always be carried out in the dark.
3. **Use of a standard when testing waste waters (method 8902):** The activated sludge must be sufficiently active. This is best carried out by using a nitrite solution (1000 mg/L) as standard inhibitor. **3.0 mL** of the supernatant and **1.0 mL** of nitrite solution are used instead of the sample. The inhibition of the dehydrogenase activity of this test sample should be between  $50 \pm 20\%$ .
4. The dry sludge mass DSM is determined by being dried at 105 °C.







**Method 8903: Evaluation of the degree of stabilization of sludge (Screening method)**

Simple, visual process to determine the degree of stabilization of sludge (activated sludge, digested sludge) upon sewage treatment plants. The greater the progression of aerobic stabilization of sludge the more inactive the activated sludge becomes or the more progressive the putrefactiveness. This is accompanied by a decline in the conversion of TTC into red triphenylformazane.

**Procedure:**

Ciclo de trabajo	
1.	<b>Preparation of samples:</b> a) Determine the dry sludge mass DSM by drying at 105 °C or estimate it $\pm 20\%$ . b) The sludge which must be investigated will be diluted with sewage treatment plant water (supernatant of sedimented activated sludge) to a dry substance content level of approx. 1 g/L. <b>Example:</b> Dry sludge mass 5 g/L → target concentration: 1 g/L → 1 + 4 dilution: 1 part sludge + 4 parts sewage treatment plant water (e.g. 4 mL sludge + 16 mL sewage treatment plant water)
2.	Add <b>1.3 mL</b> reagent <b>R1</b> into an empty test tube and subsequently fills the test tube <b>up to the brim</b> and <b>without air bubbles</b> with the prepared test solution. Seal the test tube and mix it by shaking.
3.	Incubation <b>in the dark at room temperature</b> .
4.	Visually inspect the test tube after 30, 45 and 60 min for the evidence of red colouring.
5.	<b>Evaluation:</b> If <b>no red colouring</b> can be seen after <b>60 min</b> then it is generally a predominantly stabilized sludge which has reached " <b>technical aerobic stabilization boundary</b> ". In the case of insufficiently stabilized sludge which is very putrefactive then a clearly recognizable red colouring can be observed after just 30 min but at the latest after 60 min.

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