S-YESMD

Biological measurement system for the detection of estrogenic activity in water

The **S-YES**^{MD} (*Saccharomyces*-Yeast Estrogen Screen or Speed-Yeast Estrogen Screen) is a biological measurement system for the detection of summary estrogenic activity in aqueous samples. The test is capable in particular for screening purposes of surface water and extracts. Due to the standardized production and the ready to use biosensor the working time in your lab will be reduced significantly. Within approx. 7 to 20 hours you get a result of the estrogenic effects expressed as EEQ (17β -Estradiol equivalent concentration) of your analyzed samples. The whole test can be performed under unsterile conditions.

MEASUREMENT PRINCIPLE

The test **S-YES**^{MD} uses genetic modified *Saccharomyces cerevisiae* BJ3505 yeast cells [1, 2]. Every yeast cell contains two plasmids: The receptor plasmid contains specific regulation sequences and the gene for the human estrogen receptor α . The reporter plasmid contains specific regulation sequences as well and the gene for the reporter enzyme β -Galactosidase. The binding of estrogen active substances to the receptor will subsequently activate the production of the reporter enzyme β -Galactosidase. The amount of the reporter enzyme correlates with the total concentration of estrogenic active substances in the sample. After addition of chromogenic substrate, the reporter enzyme activity can be measured photometrically. 17β -Estradiol (E2) is used as reference standard for the calibration.

[1] McDonnell et. al, 1991. In situ distinction between steroid receptor binding and transactivation at a target gene.

[2] McDonnell et. al, 1991. High level expression of biologically active estrogen receptor in Saccharomyces cerevisiae.



▲ S-YES^{MD} test kit

B-Galactosidase

HO

CPRG

Chlorophenol red

Galactose

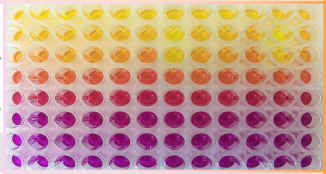
 \blacktriangle Schematic reaction of $\beta\text{-galactosidase}.$ Cleavage of CPRG into chlorophenol red and galactose

ADVANTAGES OF THE S-YES™

- Short processing time (Speed version approx. 7 hours)
- Easy handling
- No precultures necessary
- No sterile working conditions required

APPLICATIONS

- Aqueous extracts
- Drinking and mineral water
- Ground and surface water
- Process water
- Wastewater



LABORATORY REQUIREMENTS

- BSL1 laboratory
- Multichannel pipette (nominal vol. 100 μl)
- Temperature controlled shaker (T = 86 °F, Orbit 3 4.5 mm)
- Microliter centrifuge
- Photometer for microtiter plates (λ = 580 and 600 or 620 nm)



S-YESMD

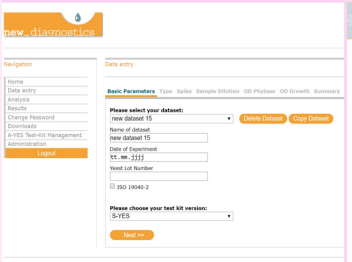
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Duration of Assay	ca. 7 – 20 h
Number of samples	max. 80
Validation	in house study
Calibration Range	0 – 400 ng/L 17β-Estradiol (E2)
Limit of detection	3.8 ng/L 17β-Estradiol (E2)

BioVAL® - SOFTWARE FOR EXPERIMENTAL DESIGN AND STATISTICAL ANALYSIS



We will give you access to BioVAL® for an easy, reliable, and uniform statistical analysis. The web-based software enables you to analyse your data in a standardized manner without special statistical knowledge. The results are presented in a comprehensive report.





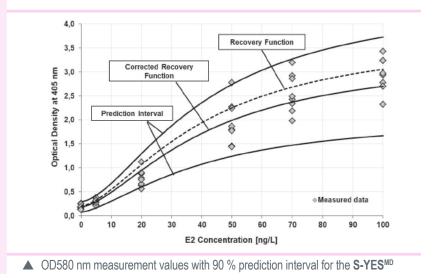
▲ Data analysis via BioVAL® webinterface

▲ Excerpt of the certificate of analysis

QuoData CERTIFICATE

The **S-YES**^{MD} test kit has been awarded the QuoData certificate of matrix comprehensive validation. This guarantees continuously high quality and reliability of our test kits.





The validation of the **S-YES**^{MD} was performed according a factorial in house Validation study with eight different water samples each spiked with different concentrations of E2.

The range of water types comprised a heterogeneous set of samples with different samples matrices e. g. surface water and methanolic extracts. The planning and evaluation of the validation was realized by QuoData GmbH.

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