

Anabolic Lab™

Analytical Lab Testing for
Anabolic Steroid Harm Reduction

<https://anaboliclab.com>

Lab Report – March 27, 2018

SIS Laboratories (Science in Sport)
Sustanon 250 (250 mg/mL)
Testosterone: Propionate, Phenylpropionate,
Isocaproate, Decanoate



If you like the attached report, please visit [our website](https://anaboliclab.com) to find out how you can support independent and objective lab testing.

Test report

Number **C 67067**

Object of analysis: SIS Sustanon 250

Customer: AnabolicLab.com **Samples:** 1

Batch: 001-30540, Mfg. 01/2016, Exp. 01/2020 **Receipt:** 28.02.2018


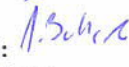
Subject: Content, Identification

Method(s): HPLC **Ansatz-datum:** 09.03.2018

Sample / Analysis:	Method	Status	Result	Specification	complies
<u>Testosterone Propionate:</u>					
Identification	HPLC	N	complies	complies	yes
Content	HPLC	N	53.7 mg / ml	50 mg / ml	-
<u>Testosterone Phenylpropionate:</u>					
Identification	HPLC	N	complies	complies	yes
Content	HPLC	N	52.7 mg / ml	50 mg / ml	-
<u>Testosterone Isocaproate:</u>					
Identification	HPLC	N	complies	complies	yes
Content	HPLC	N	55.3 mg / ml	50 mg / ml	-
<u>Testosterone Decanoate:</u>					
Identification	HPLC	N	complies	complies	yes
Content	HPLC	N	102.9 mg / ml	100 mg / ml	-
TAMC	SOP M 006	N	< 1 KBE/ml	-	-
TYMC	SOP M 006	N	< 1 KBE/ml	-	-

Method-status: G = GMP A = accredited V = generally validated P = validated on product N = not validated
 E = external lab

Remarks: The sample(s) mentioned above have been analysed as they have been sent to us by the client. We have not controlled external processes like manufacturing, labelling, sampling, shipping, storage. These results are for information only and do not compensate for correct quality control by the manufacturer/distributor. Generally, pharmaceutical products have to be produced and distributed under full GMP/GDP regime, including GMP-compliant analyses with validated methods. This report cannot be used for commercial reasons incl. product release and/or quality control. It is not allowed to use this analysis in the context of doping/sports/competitions or any other illegal action, neither by the athlete nor by the tutor/trainer or any other person.

Signatures: controlled & released:  **Staff:**  **completed:** 27.03.2018

The enclosed results refer exclusively to the object of examination described above. The measuring accuracy is available on request. Only handwritten signatures are valid.
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