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# Qant-iT<sup>™</sup> PicoGreen<sup>®</sup> dsDNA Quantification V.2

In 1 collection

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#### External link: https://www.thermofisher.com/order/catalog/product/P11496

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### Protocol status: Working We use this protocol and it's working

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# Abstract

The following protocol is intended for the quantification of double-stranded DNA using <u>Quant-iT<sup>™</sup>PicoGreen<sup>®</sup></u> <u>dsDNA Assay Kit (ThermoFisher</u>). This protocol is a simplified and condensed version of the <u>full protocol</u> from the manufacturer. The procedure described here is for 96 reactions. If samples are run in duplicates, then this should allow quantifying 40 samples.

## Attachments



240KB

# Materials

#### MATERIALS

🛿 🔀 Quant-iT™ PicoGreen™ dsDNA Assay Kit Invitrogen - Thermo Fisher Catalog #P11496

### STEP MATERIALS

X Quant-iT<sup>™</sup> PicoGreen<sup>™</sup> dsDNA Assay Kit Invitrogen - Thermo Fisher Catalog #P11496

### **Protocol materials**

X Quant-iT<sup>™</sup> PicoGreen<sup>™</sup> dsDNA Assay Kit Invitrogen - Thermo Fisher Catalog #P11496

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## Safety warnings

Quant-iT<sup>™</sup> PicoGreen<sup>®</sup> reagent is classified as Not Hazardous. Nevertheless, the user should always consult the MSDS accompanying any of the reagents and apparatus described in this protocol.

# Before start

- 1. This protocol is optimised for measuring an entire 96-well plate. It assumes that 16 wells will be used for measuring the standards and the blank samples (in duplicates) and 80 wells will be used for measuring unknown DNA samples (typically in duplicates).
- 2. The protocol can be easily adjusted for a lower number of samples by reducing the volume of the working solutions of the reagents. Note though that enough TE should be retained for diluting the standard stock solution (490 or 680  $\mu$ l), for potentially diluting the unknown samples, if their concentration is too high, and for accounting for pipetting errors. To fill the plate, 19.2 ml of TE is needed. So if only 40 wells are to be used for measuring unknown samples prepare about  $50/96 \times 22 \approx 11.5$  ml of TE buffer.
- 3. The dynamic range of the assay is between 50 pg ml<sup>-1</sup> to 1000 ng ml<sup>-1</sup>. This translates into DNA sample concentrations of 0.05-5 ng  $\mu$ l<sup>-1</sup> and 1-200 ng  $\mu$ l<sup>-1</sup> in the low-range and high-range assays, respectively. Samples with higher DNA concentration need to be diluted (e.g. in DNase-free water or TE buffer).
- 4. Note that some compounds that can be present as DNA contaminations (e.g. salts, ethanol, detergents, proteins) are claimed by the manufacturer to not interfere with the measurement. Please refer to the full protocol for a list of these compounds and their effect on the measurement. Also, equimolar presence of ssDNA and RNA in the sample should have only minimal effect on the quantitation results.

Pre	pare reaction	53m
1	Take out all reagents from the fridge and bring them to room temperature. Take out the DNA samples from the freezer. DNA samples should be slowly thawed on ice.	20m
	Note	
	Quant-iT <sup>™</sup> PicoGreen <sup>®</sup> dsDNA reagent is dissolved in dimethylsulfoxide (DMSO), which freezes below 19 °C. The reagent must be completely thawed before using it by bringing it to room temperature. After the reagent thawed, it is advisable to briefly vortex the tube to make sure it is adequately mixed and to spin it down in a centrifuge or a mini centrifuge.	
	Note	
	Quant-iT™PicoGreen <sup>®</sup> dsDNA reagent is light sensitive and should be protected from light at all times.	
	Want-iT™ PicoGreen™ dsDNA Assay Kit <b>Invitrogen - Thermo</b> Fisher Catalog #P11496	
_		
2	Prepare 22 ml 1X TE buffer by pipetting 1.1 ml of 20X TE buffer into 20.9 ml of nuclease- free water into a sterile and nuclease-free 50 ml tube. Mix by inverting the tube several times.	2m
	A 1.1 mL 20X TE buffer	
	▲ 20.0 mL nuclease-free water	
3	<b>For high-range quantification:</b> Dilute the DNA-standard stock solution (λ DNA 100 ng $\mu$ l <sup>-1</sup> ) to a final concentration of 2 ng μl <sup>-1</sup> by mixing 10 μl λ DNA-standard stock solution with 490 μl 1X TE buffer.	2m
	$\Delta$ 10 µL $\lambda$ DNA-standard stock solution	
	490 μL 1X TE buffer	

Prepare a 40-fold dilution of the 2 ng  $\mu$ l<sup>-1</sup> DNA-standard work solution by mixing 5  $\mu$ l of the 2 ng  $\mu$ l<sup>-1</sup> DNA-standard work solution with 195  $\mu$ l 1X TE buffer to yield a 0.05 ng  $\mu$ l<sup>-1</sup> DNA-standard work solution.

- $\triangleq$  5 µL diluted DNA-standard solution
- 👗 195 μL 1X TE buffer
- 4 If needed, prepare a dilution of each sample in 1X TE buffer so that the reading will be within the dynamic range.

Note

It is advisable to run samples in duplicates for a more accurate quantification

5 Prepare PicoGreen<sup>®</sup> work solution: 9950 μL 1X TE buffer + 50 μL PicoGreen<sup>®</sup> into a sterile and nucleic-acids free 50 ml tube. Mix and protect from light.

🛓 9950 μL 1X TE buffer

🕹 50 μL PicoGreen®

6 Prepare the following standard mixture in the first two columns of the black, sterile, 96well plate:

	Assay version	Dilut ed DNA std. (µl)	1X TE buffe r (μl)	Final DNA amou nt (ng)
	High-range (1-200 ng µl-1)	100	0	200
	Use 2 ng µl-1 standard	50	50	100
		10	90	20
		1	99	1
		0	100	0
	Low-range (50 pg µl-1 - 5 ng µl-1)	100	0	5
	Use 0.05 ng μl-1 standard	50	50	2.5
Γ		10	90	0.5
		1	99	0.05
		0	100	0

2m

10m

Equipment	
96-well microtiter plate	NAME
Nunc	BRAND
265301	SKU
https://www.thermofisher.com/order/catalog/product/152028#/152028	
black, flat bottom	SPECIFICATIONS

7 Pipette 99  $\mu$ l of 1X TE buffer in the remaining wells.

4 99 μL 1X TE buffer

#### Note

Tip: use a mechanical or electronic dispenser during this step and step no. 9 to speed up the work.

	•		and the
Equ	lip.	me	ent

Multipette E3	NAME
Eppendorf	BRAND
4987000010	SKU
http://multipette-system.eppendorf.com/	LINK
electronic dispenser	SPECIFICATIONS

8 Pipette 1  $\mu$ l of the unknown DNA samples in the remaining wells.

	- 1 μL of DNA san	nple		10m
	Note			
			ne concentration is expected to be higher than the in case the concentration is expected to be below the	
9	Pipette 100 μL of the unknown sample we		rk solution in each well, including the standard and	2m
	Δ 100 µL PicoGree	en work solution		
10	Protect the 96-well	plate from light ar	nd incubate for 2-5 min at room temperature.	5m
Меа	asure samples			5m
11	Place the plate in a parameters:	plate reader and r	measure the fluorescence according to the following	5m
	Excitation	~480 nm		Ŧ
	Emission	~520 nm		
	Integration time	40 s		
	Lag time	0 s		
	Gain	Optimal		
	Number of flashes	10		

#### Note

Shaking

Calculated well

It is also possible to set the gain to a fixed value (e.g. 100). If the fluorescence values of the standard drop over time this could indicate damage to the reagents or the DNA standard.

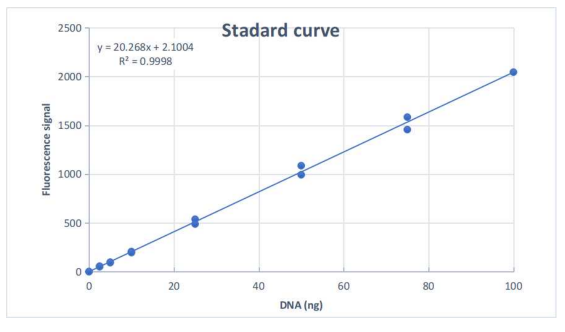
highest standard

5 s

Equipment	
Synergy 2	NAME
absorbance microplate reader	TYPE
BioTek	BRAND
Synergy2	SKU
https://www.biotek.com/products/detection/	LINK

12 Plot the measured fluorescent values of the standard samples against their known concentrations and fit a linear curve using linear regression. Make sure that the coefficient of determination (R2) is close to 1 (typically > 0.99). Calculate the DNA concentrations in the unknown samples using the slope and intercept parameters of the linear equation. Output values you obtained are in ng  $\mu$ <sup>-1</sup>, assuming 1  $\mu$ l of each sample was used.

See attached example spreadsheet.



10m

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#### Note

Do not forget to account for any dilutions when calculating the concentration of the DNA in the unknown samples.

PicoGreen\_example.xlsx