



Nickel Columnation August And

(Catalog #K510-100; 100 assays; Store at room temperature)

I. Introduction:

Nickel is one of four ferromagnetic elements (symbol Ni, at. Num. 28). Several enzymes depend on nickel for activity, including some ureases, carbon monoxide dehydrogenases (methane forming enzymes which reduce CO_2 to CH₄) and some hydrogenases which allow the production or removal of H₂. Most of these activities are found in the archaebacteria. Nickel forms complexes with sulfhydryl compounds with significant absorbance in the UV/visible region in the presence of other ions. BioVision's Nickel Assay kit provides a simple method of quantifying Ni²⁺ in a variety of samples. The assay takes advantage of reaction of Ni²⁺ with mercaptoethanol in borate buffer to form a complex with strong absorbance bands from ~300 to 600 nm. Fe²⁺ and Co^{2+} interfere with the assay, therefore extra steps (as described below) must be taken to subtract the interference in order to determine the correct Nickel concentration in mixed samples. Other ions tested (Mn²⁺, Cu²⁺, Zn²⁺) do not interfere with the assay, presumably no other ionic species are present in high enough concentration to interfere with the reaction. The assay is a simple method of quantifying Ni²⁺ in a variety of samples, which gives a linear range of 2 to 50 nmol Nickel containing less than 25 nmol Cobalt.

II. Kit Contents:

Components	K510-100	Cap Code	Part Number
Nickel Assay Buffer	20 ml	WM	K510 -100-1
Nickel Reagent	1 ml	Green	K510 -100-2
Nickel Chloride Standard (1.0 µmol)	Iyophilized	Yellow	K510 -100-3

III. Reagent Preparation and Storage Conditions:

Store kit at room temperature, keep tightly capped.

Nickel Assay Buffer and Nickel Reagent: Ready to use as supplied. Store at room temperature. Stable for at least 6 months.

Nickel Standard: Dissolve in 1 ml dH₂O to make a 1 mM solution. Store at room temperature.

IV. Nickel Assay Protocol:

- Standard Curve Preparations: Add 0, 10, 20, 30, 40, 50 µl of the Nickel standard to a series of wells. Adjust volume to 200 µl/well with Nickel Assay Buffer to generate 0, 10, 20, 30, 40 and 50 nmol per well of the Nickel Standard.
- **2. Sample Preparation:** Sample Nickel concentrations can vary over a wide range. Take samples between 10-100 μl and adjust volume to 200 μl with Nickel Assay Buffer for each well. For unknown samples, different sample amounts should be tested to ensure the readings are within the standard curve linear range.

Note: In the absence of Fe^{2*} and Co^{2*} in samples, the protocol requires reading OD405 nm only. In the presence of Fe^{2*} and Co^{2*} in samples, the protocol requires two separate readings at two different wavelengths to correct interference from Fe^{2*} and/or Co^{2*} .

- Reading 1: Read OD of the samples and standards at 330 nm and 405 nm before adding the Nickel Reagent. This OD is due only to Fe²⁺and reagent background. Call these measurements OD330₁ and OD405₁.
- **4. Development:** Add 10 µl of Nickel Reagent to all standard and sample wells. Incubate at room temperature for 30 min to form complex.
- 5. Reading 2: Read OD at 330 nm and 405 nm. Call these measurements OD330₂ and OD405₂.
- 6. Nickel Determination in the absence of Fe²⁺, or/and Co²⁺: Subtract reading 1 (OD405₁) from reading 2 (OD405₂) to get the corrected reading ΔOD405. Plot the standard curve. Apply corrected ΔOD405 of unknown samples to the standard curve to determine Ni²⁺ amount in the reaction wells (Ay). Calculate Nickel concentration as in step 8 (without the step 7 corrections).

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- **7.** Nickel Determination in the presence of Fe²⁺ or/and Co²⁺: Subtract the 0 Nickel OD reading from all standard, sample readings to correct absorbance due to buffer and plate.
 - 1) Remove interference at 330 nm due to Fe^{2+} : In the absence of Nickel Reagent, OD330₁ is contributed only by Fe^{2+} . After adding Nickel Reagent, Fe^{2+} contribution to the OD330₂ can be calculated as follows: $FeOD330_2 = 1.82 \times OD330_1$. Subtract the FeOD330₂ value from total OD330₂ to get corrected OD330, Δ FeOD330₂ = OD330₂ FeOD330₂, which is contributed by Ni²⁺ and Co²⁺.
 - 2) Remove interference at 405 nm due to Fe: In the absence of Nickel Reagent OD405₁ is contributed only by Fe²⁺. After adding Nickel Reagent, Fe²⁺ contribution to OD405₂ can be calculated as follows: FeOD405₂ = 1.65 X OD405₁. Subtract FeOD405₂ value from total OD405₂ reading, to get corrected OD405₂ reading, ΔFeOD405₂ = OD405₂ FeOD405₂, which is contributed by only Ni²⁺ and Co²⁺.
 - 3) Remove interference due to Co^{2^+} : Calculate the ratio of $\Delta FeOD330_2$ and $\Delta FeOD405_2$: $\Delta FeOD330_2/\Delta FeOD405_2$. The ratio should fall between 0.925 (100% Co) and 2.8125 (100% Ni). Subtract 0.925 from the ratio calculated and divide that result by 1.8875, ($\Delta FeOD330_2/\Delta FeOD405_2 - 0.925$)/1.8875 is the percentage of absorbance due to Ni²⁺. Multiply that percentage by $\Delta FeOD330_2$ to get Nickel absorbance at OD330₂, $\Delta FeCOD330_2$ in samples.
- 8. Calculation: Plot the standard curve (△OD405, or △OD330). Calculate sample Nickel reading △OD405 from step 6 for samples without Fe²⁺ and Co²⁺, or △FeCoOD330₂ from step 7 for samples with Fe²⁺ or/and Co²⁺ interference. Apply the sample readings to the standard curve to get Ni²⁺ amounts (Ay) in the reaction well.

Nickel Concentration = Ay/Sv (nmol/ml)

Where: Ay is the amount of Ni²⁺ (nmol) in your sample from the standard curve. Sv is the sample volume (in ml) added to the sample well. Nickel molecular weight: 58.7 g/mol.



Nickel Standard Curve: Assays were performed following the kit protocol.

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RELATED PRODUCTS:

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Phosphate Assay Kit	
Magnesium Assay Kit	
Urea Assay Kit	
Glutathione Assay Kit	

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GENERAL TROUBLESHOOTING GUIDE:

Problems	Cause	Solution				
Assay not working	Use of ice-cold assay buffer	Assay buffer must be at room temperature				
	Omission of a step in the protocol	Refer and follow the data sheet precisely				
	Plate read at incorrect wavelength	Check the wavelength in the data sheet and the filter settings of the instrument				
	Use of a different 96-well plate	Fluorescence: Black plates (clear bottoms) ; Luminescence: White plates ; Colorimeters: Clear plates				
Samples with erratic readings	Use of an incompatible sample type	Refer data sheet for details about incompatible samples				
	Samples prepared in a different buffer	Use the assay buffer provided in the kit or refer data sheet for instructions				
	Samples were not deproteinized (if indicated in datasheet)	Use the 10 kDa spin cut-off filter or PCA precipitation as indicated				
	Cell/ tissue samples were not completely homogenized	Use Dounce homogenizer (increase the number of strokes); observe for lysis under microscope				
	Samples used after multiple free-thaw cycles	Aliquot and freeze samples if needed to use multiple times				
	Presence of interfering substance in the sample (e.g. metal ion chelators)	Troubleshoot if needed, deproteinize samples				
	Use of old or inappropriately stored samples	Use fresh samples or store at correct temperatures till use				
Lower/ Higher readings in Samples and Standards	Improperly thawed components	Thaw all components completely and mix gently before use				
	Use of expired kit or improperly stored reagents	Always check the expiry date and store the components appropriately				
	Allowing the reagents to sit for extended times on ice	Always thaw and prepare fresh reaction mix before use				
	Incorrect incubation times or temperatures	Refer datasheet & verify correct incubation times and temperatures				
	Incorrect volumes used	Use calibrated pipettes and aliquot correctly				
Readings do not follow a linear pattern for Standard curve	Use of partially thawed components	Thaw and resuspend all components before preparing the reaction mix				
	Pipetting errors in the standard	Avoid pipetting small volumes				
	Pipetting errors in the reaction mix	Prepare a master reaction mix whenever possible				
	Air bubbles formed in well	Pipette gently against the wall of the tubes				
	Standard stock is at an incorrect concentration	Always refer the dilutions in the data sheet				
	Calculation errors	• Recheck calculations after referring the data sheet; Correct for Fe ²⁺ and Co ²⁺				
	Substituting reagents from older kits/ lots	Use fresh components from the same kit				
Unanticipated results	Measured at incorrect wavelength	Check the equipment and the filter setting				
	Samples contain interfering substances	Troubleshoot if it interferes with the kit				
	Use of incompatible sample type	Refer data sheet to check if sample is compatible with the kit or optimization is needed				
	Sample readings above/below the linear range	Concentrate/ Dilute sample so as to be in the linear range				
Note: The most probable list of caus	lote: The most probable list of causes is under each problem section. Causes/ Solutions may overlap with other problems.					