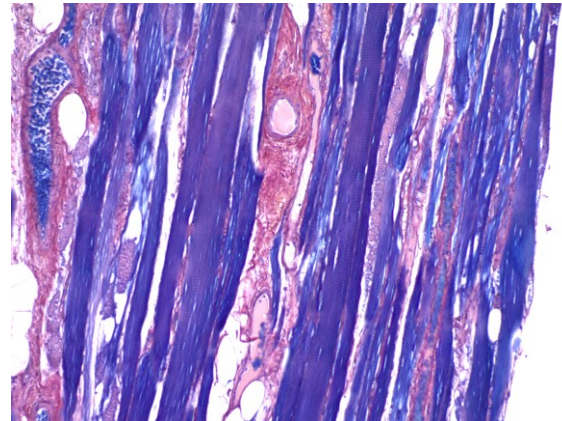


PTAH Stain Kit (Phosphotungstic Acid Hematoxylin)

Description: The PTAH Stain Kit is intended for use in the histological visualization of collagen, striated muscle, glial fibers and collagen without using Zenker’s Fixative with Mercuric Chloride as a mordant. This kit may be used on formalin-fixed, paraffin-embedded sections.

Fibrin, Striated Muscle, Glial Fibers:	Blue to Purple
Collagen:	Light Orange/Salmon to Brownish/Red
Nuclei:	Blue to Purple

Uses/Limitations: Not to be taken internally.
For In-Vitro Diagnostic use only.
Histological applications.
Do not use if reagents become cloudy.
Do not use past expiration date.
Use caution when handling reagents.
Non-Sterile.



Control Tissue: Striated Muscle


Ordering information regarding individual components on back page!

Kit Contents:

<u>Item #</u>	<u>Components</u>	<u>Volume</u>	<u>Storage</u>
ZCS500	Zinc Chloride Solution (10%)	500 ml	18-25° C
FAS125	Ferric Ammonium Sulfate	125 ml	18-25° C
HPA125	PTAH Solution	125 ml	18-25° C

Precautions: Avoid contact with skin and eyes.
Harmful if swallowed.
Follow all Federal, State, and local regulations regarding disposal.

Storage: 18° C  25° C

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Logan, UT 84321
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CE IVD

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Procedure (60° C. Water Bath):


1. Deparaffinize sections if necessary and hydrate to distilled water.
 2. Pour Zinc Chloride Solution (10%) into plastic staining jar and set in 60° C. water bath for 10 minutes to equilibrate temperature.
 3. Place slide in warmed Zinc Chloride Solution (10%) and incubate for 20 minutes at 60°C.
 4. During step 3, pour Ferric Ammonium Sulfate Aqueous Solution into a second plastic staining jar and set in 60° C. water bath for 10 minutes to equilibrate temperature.
 5. Rinse slide in running tap water for 1 minute.
 6. Rinse in distilled water for 1 minute.
 7. Place slide in warmed Ferric Ammonium Sulfate Aqueous Solution and incubate for 5 minutes at 60°C.
 8. During step 7, pour Phosphotungstic Acid Hematoxylin Solution into a third plastic staining jar and set in 60° C. water bath for 10 minutes to equilibrate temperature.
 9. Rinse slide in running tap water for 2 minutes.
 10. Rinse in distilled water for 1 minute.
 11. Place slide in warmed Phosphotungstic Acid Hematoxylin Solution and incubate for 60 minutes at 60°C.
 12. Differentiate section in 95% Reagent Alcohol. Check section using microscope for proper differentiation.
- Note: Graded alcohols will remove some stain.
13. Dehydrate in 3 changes of Absolute Alcohol.
 14. Clear in 3 changes of fresh Xylene or Xylene Substitute, and mount in synthetic resin.

Procedure (Microwave):

Equipment Needed: 500 Watt Microwave Oven

1. Deparaffinize sections if necessary and hydrate to distilled water.
2. Place slide to fresh distilled water for 1 minute.
3. Pour 50 ml of (cat# ZCS) Zinc Chloride Solution (10%) into plastic coplin jar and heat in microwave for 20 seconds on high power. Remove jar and stir solution to equalize temperature. Return coplin jar to microwave and heat for 10 seconds on high power. Remove jar and stir solution to equalize temperature.
4. Place slide in coplin jar and incubate for 15 minutes.
5. Rinse slide in running tap water for 1 minute.
6. Rinse in distilled water for 1 minute.
7. Place slide in 25 ml (cat# FAS) Ferric Ammonium Sulfate Aqueous Solution, heat in microwave for 15 seconds on high power and incubate for 2 minutes.
8. Rinse slide in running tap water for 2 minutes.
9. Rinse in distilled water for 1 minute.
10. Heat 25 ml (cat# HPA) Phosphotungstic Acid Hematoxylin Solution in microwave for 20 seconds on high power. Remove and agitate to equalize temperature of solution. Place slide in stain, agitate and incubate for 15 minutes. Reheat solution for 10 seconds on high power, agitate and incubate for another 15 minutes.
11. Differentiate section in 95% Reagent Alcohol. Check section using microscope for proper differentiation.
12. Dehydrate in 3 changes of Absolute Alcohol.
13. Clear in 3 changes of fresh Xylene or Xylene Substitute, and mount in synthetic resin.

Storage: 18° C  25° C

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
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
References:

- Shapiro, S.H., Sohn, L.C.; Rapid Microwave Phosphotungstic Acid-Hematoxylin Stain for Paraffin and Glycol Methacrylate Sections; The Journal of Histotechnology; Volume 17, Number 2, June 1994, pages 125-126.

Bulk Reagent Ordering Information and Current Pricing at www.scytek.com

Description:	Catalog #	Volume
Zinc Chloride Solution (10%)	ZCS500	500 ml
	ZCS999	1000 ml
Ferric Ammonium Sulfate Aqueous Solution	FAS125	125 ml
	FAS500	500 ml
	FAS999	1000 ml
Phosphotungstic Acid Hematoxylin Solution	HPA125	125 ml
	HPA500	500 ml
	HPA999	1000 ml

 Storage: 18° C  25° C



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