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#### TECHNICAL DATA SHEET 787

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# **Preparing Colloidal Gold for Electron Microscopy**

#### **Preparation of Colloidal Gold Conjugates**

Colloidal gold has been used for centuries in the preparation of stained glass for windows and fine glassware. In recent years, colloidal gold particles have become a useful tool for microscopists. Colloidal gold particles are especially useful for biological electron microscopy. Some of the reasons why are listed below.

- Homogeneous preparations of particles varying in size from 3 nm to 20 nm can be easily prepared.
- Colloidal gold suspensions are inexpensive to prepare.
- Most proteins can be easily coupled to colloidal gold particles.
- Proteins coupled to gold particles do not appear to lose their biological activity.
- The colloidal gold particles can be easily seen in the electron microscope.
- Colloidal gold probes can be used for light microscopy. The larger gold particles can be directly observed by the light microscope. Smaller particles are detected by silver enhancement or epipolarized illumination.
- The same probes can be used for both LM and TEM immunocytochemistry.

#### Preparation procedure for producing gold sols

This page will explain a simple, published protocol for preparing colloidal gold particles, how to couple these particles to protein A and how to purify the probes after they have been made.

To make 100 ml of gold sol, two stock solutions have to be prepared.

#### Solution A

80 ml distilled water and 1 ml 1% aqueous gold chloride.

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4 ml 1% tri-sodium citrate. 2H2O + 16 ml H2O + variable amount of 1% tannic acid (Cat.#04459). (See table 1)

When 1 ml or more tannic acid is needed, add an equal amount of 25mM potassium carbonate for pH adjustment.

Warm up solutions A and B to 60°C and mix them while stirring. When the red color has formed heat up to 95°C and cool the solution on ice. The larger particles (where lower concentrations of tannic acid are used) take longer to form and the red color can take up to one hour to develop.

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## **Coupling the Protein A to the Gold Particles**

A gold sol will bind proteins more efficiently when the pH of the solution is close to the pI of the protein (for protein A the pI is pH 5.1). A tighter binding occurs at higher pH but this may have a denaturing effect on the protein making the probe less effective. Having too much protein coupled to the gold particles may also be disadvantageous: some of the weakly bound protein may detach from the particles. This will make the probe less effective because the free protein will compete for binding sites with the gold-labelled protein. Horrisberger and Clerc (1985, Labelling of colloidal gold with protein A. A quantitative study. Histochemistry, 82, 219-223) recommend binding the protein A to colloidal gold at pH 6.0.

Check the pH of the gold sol with pH paper (the gold sol will block a pH electrode) and adjust the pH with 0.1N sodium hydroxide.

Protein A (Boehringer Mannheim) is dissolved in distilled water at 1mg/ml. A microtitration assay will show the correct amount of protein A to add to the gold sol (between 4-6µg/ml).

Add the protein A while stirring the gold sol. After 5 min add 10% bovine serum albumin (BSA) in PBS to a final concentration of 0.2% (2ml/100ml) to maximally stabilize the sol.

#### **Purification of the Protein A-gold**

The protein A-gold is centrifuged in a Ti 70 rotor in a Beckman ultracentrifuge at the appropriate speed (see table 2) for 30 min at 4°C. At the correct speed the gold particles will settle to the bottom of the tube as a loose pellet. Remove the supernatant without disturbing the pellet and re-suspend the loose part of the pellet in PBS containing 0.2% BSA.

A second centrifugation step down a gradient will remove any gold particles of the wrong size. This is done on a 10-30% continuous glycerol gradient at 4°C. Layer 1-2 ml of protein A gold onto the top of the gradient, spin in an SW40 rotor at the appropriate speed (see table 3) for 45 min. at 4°C. The dark red band in the middle of the gradient is collected. All aggregated protein A gold particles will have been removed.

#### **Storage**

Immunogold probes may lose activity within weeks, due to the dissociation of the proteins from the gold particles.

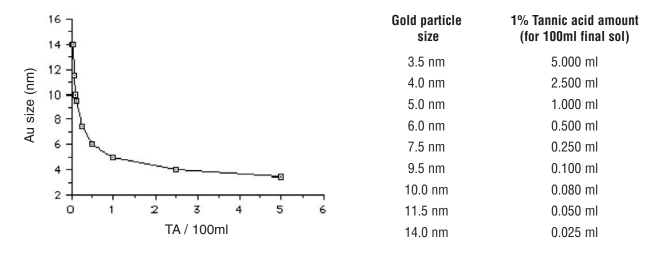
Dialyse the protein A gold against 50% glycerol in PBS and store at -20°C or freeze down small aliquots in liquid nitrogen and store at -70°C.

To determine the concentration at which to use the protein A gold, measure the optical density (OD) at 520 nm of a 1:100 diluted solution in PBS. Use a dilution with an O.D. of between 0.05 and 0.1 where there is no significant background. If using a primary antibody then the optimal dilution of this first antibody must be known. Sections can be treated with protein A gold alone to determine the background labeling.

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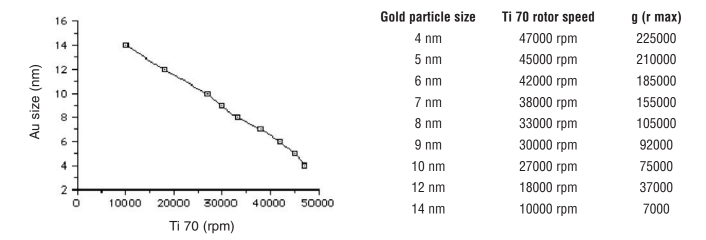
**2** 

**Table 1:**The influence of the tannic acid concentration, during gold sol formation, on the size of the gold particles.



When 1 ml or more of tannic acid is used an equal amount of 25 mM potassium carbonate must be added to neutralize solution B.

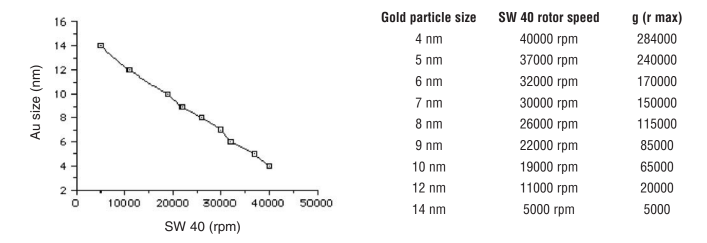
**Table 2:** First centrifugation, using a Ti 70 rotor for 30 min at 4°C. This step will concentrate the colloidal gold probe in the bottom of the tube.



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**Table 3:**Second centrifugation using SW 40 rotor for 45 min at 4°C. This will separate the different sizes of gold particle along the gradient.



The 4 nm gold must be centrifuged for 1 hr for best results. The 5nm gold can also be centrifuged at 35000 rpm for 1 hr.

### Coupling bovine serum albumin (BSA) to gold particles

BSA-gold is a useful marker for studying the endocytic processes in mammalian cells. Typically, the living cells are incubated in a BSA-gold suspension with a final OD, at 520 nm, of 5, meaning that a large amount of gold probe is needed for these experiments. The above protocol for preparing protein A-gold can be followed for preparing BSA-gold but BSA is substituted for the protein A (we use  $2.4 \,\mu g/ml$ ). The centrifugation values are the same.

After the gold probe has been concentrated and purified it is dialized against PBS, or culture medium, before being added to the living cells.

#### References

Slot, J. W. & H. J. Geuze 1981. Sizing of protein A-colliodal gold probes for immunoelectron microscopy. J. Cell Biol. 90, 533-536. Slot, J. W. & H. J. Geuze 1985. A new method of preparing gold probes for multiple-labeling cytochemistry. Europ. J. Cell Biol. 38, 87-93.

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## **Ordering Information**

Catalog #	Description	Size
09285-50	Colloidal Gold Solution, 0.005%	50ml

#### **Adjuncts/Additional Products**

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Catalog #	Description	Size	
04459-100	Tannic Acid, EM Grade	100g	
00740 44	Large Cold labeling grounds for (TEM). He against d (OO) Form	1001	
22716-11	ImmunoGold labeling reagents for (TEM) - Unconjugated (GC), 5nm	100ml 100ml	
22717-11 22718-100	ImmunoGold labeling reagents for (TEM) - Unconjugated (GC), 10nm	100ml	
22719-100	ImmunoGold labeling reagents for (TEM) - Unconjugated (GC), 15nm ImmunoGold labeling reagents for (TEM) - Unconjugated (GC), 20nm	100ml	
22720-100	ImmunoGold labeling reagents for (TEM) - Unconjugated (GC), 40nm	100ml	
22703-100	ImmunoGold labeling reagents for (TEM) - Unconjugated (GC), 60nm	100ml	
22700 100	minulodold labeling reagents for (TEM) Onconjugated (do), comm	1001111	
24876-1.25	Amino PEGylated Gold Nanoparticles, 15nm	1.25ml	
24877-1.25	Amino PEGylated Gold Nanoparticles, 20nm	1.25ml	
24878-1.25	Amino PEGylated Gold Nanoparticles, 30nm	1.25ml	
24870-1.25	Carboxyl PEGylated Gold Nanoparticles, 15nm	1.25ml	
24871-1.25	Carboxyl PEGylated Gold Nanoparticles, 20nm	1.25ml	
24872-1.25	Carboxyl PEGylated Gold Nanoparticles, 30nm	1.25ml	
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24879-1.25	Methyl PEGylated Gold Nanoparticles, 15nm	1.25ml	
24880-1.25	Methyl PEGylated Gold Nanoparticles, 20nm	1.25ml	
24881-1.25	Methyl PEGylated Gold Nanoparticles, 30nm	1.25ml	
24873-1	Neutravidin PEGylated Gold Nanoparticles, 15nm	1ml	
24874-1	Neutravidin PEGylated Gold Nanoparticles, 20nm	1ml	
24875-1	Neutravidin PEGylated Gold Nanoparticles, 30nm	1ml	
24688-5	Mercaptyalkyl PEG Gold Nanoparticles	5ml	
24689-5	Naked Gold Nanoparticles	5ml	
24690-5	Thiol Capped Gold Nanoparticles (2-5nm)	5ml	

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