



## BIOTECH SUPPORT GROUP

### ProCipitate™

*Superior Substitute to Phenol/Chloroform for Hemoglobin & Protein Removal,  
Isolation of DNA/RNA*

- ❖ Removes protein contaminants & leaves DNA & RNA soluble and unreacted
- ❖ Ideal for applications when the alternative kits don't fit, or are not optimal
- ❖ Adaptable to any sample size and can be automated
- ❖ Pathogen and infectious disease testing
- ❖ Tissue and paraffin-embedded tissues



The **ProCipitate™** strategy is opposite to common prep strategies, as instead of binding nucleic acids, the proteins and especially - Hemoglobin, is efficiently depleted with no interaction with the soluble nucleic acids.

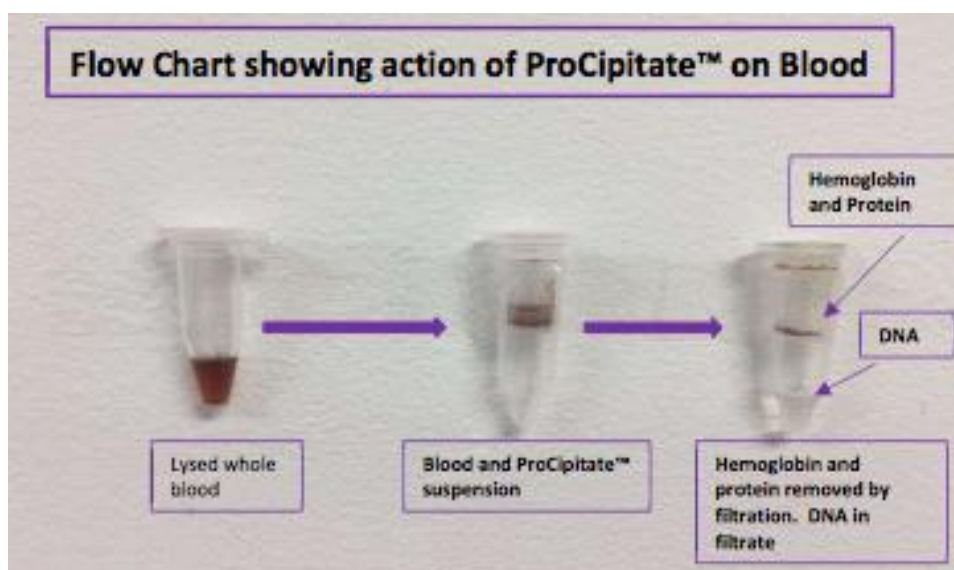
**ProCipitate** is a unique protein binding reagent developed from patented solid-phase polyelectrolytes. These elastomeric polymer suspension reagents are prepared in an extended state due to strong electrostatic repulsion of the repeating polymeric acid groups. Upon interaction with proteins, salt bridges form and the resultant complex collapses to a lower energy state, expelling water much like dehydration processes taking place within solvent precipitations. Consequently, aggregation of proteins is strongly promoted and occurs even in high ionic strength or surfactant containing solutions. Most importantly, nucleic acids remain unreacted and are quantitatively recovered in solution.

In this way, **ProCipitate™** is characteristic of phenol/chloroform separation, a long established benchmark for nucleic acid isolation. However, **ProCipitate™** is non-volatile, non-hazardous, and has the additional benefits of solid-phase suspensions; that is - the adaptability to filtration and automation. **ProCipitate™** has been on the market for over 20 years being used throughout the Human Genome Sequencing Project. It is routinely used for improving the yield consistency and protein depleted quality of DNA. Such improvements have been cited in sequence and PCR quality for a variety of applications, most notably in the template preparation of large insert plasmids (cosmids & BACs) and PCR suitability for infectious agents and from paraffin-embedded tissues.

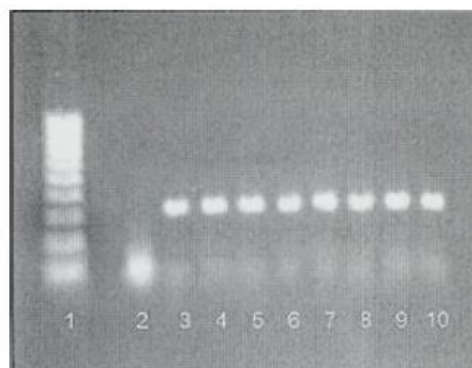
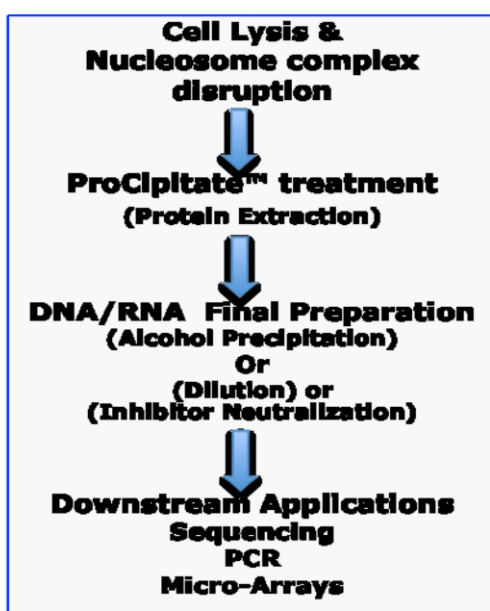
**ProCipitate™** has also been used for enrichment of other macromolecules including viruses, proteoglycans, polysaccharides, glycolipids, and highly substituted polymer conjugates (i.e., PEG), which serve to mask salt bridge formation and retain solubility. A full list of references is provided at the end of this product sheet.



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**ProCipitate™** and related products can be customized to fit specific needs. It also can be supplied in bulk quantities. Please contact our sales office or any of our worldwide distributors for more information.



Lane 1: 100-1000 base pair ladder  
Lane 2: Negative control  
Lanes 3-10: PCR amplicons (HLA-DR-Beta primers, 32 cycles) from 1 ng template DNA purified from whole blood



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### Considerations for optimal use

The optimal use of **ProCipitate™** for nucleic acid isolation should consider these 4 necessary processes:

1. The cells/tissue must be sufficiently lysed so that the intra-cellular fraction is released into the surrounding media, and the nucleosome (histone protein/ DNA complex) is efficiently disrupted.
2. The amount of **ProCipitate™** required will depend on the starting sample protein load, guidelines for which are provided below, and
3. The soluble DNA upon treatment, must be finally prepared to neutralize any inhibitory effects of the lysis condition. Typically, this is done by alcohol precipitation, dilution, or chemical neutralization; for which some of these protocols are documented in the references provided. Alternatively, DNA/RNA binding filters can be incorporated.

Because **ProCipitate™** is reactive under diverse lysis conditions, the user has great latitude in designing a protocol optimized for their own particular application and sample type.

Sample Size	ProCipitate™ Typical Usage
1 ml Yeast Culture Genomic DNA	200 µl
Mouse Tail Genomic DNA	200 µl
1 mm Plant Leaf	50 µl
2.0 ml culture BAC Preps	80 µl
5µm paraffin-embedded tissue	200 µl
250 µl culture Plasmid Preps	20 µl
200 ml Large Scale BAC Preps	5 ml
Dried Blood Card or ~ 40 µl Whole Blood	400 µl
200µl lysed cell pellet	200 µl



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**ProCipitate™** performs optimally in a final pH range of approximately 4 to 6, however the polyelectrolyte is sufficiently acidic (pH 4) to lower the final reaction pH to within its optimal working range in most applications.

Product	Size	Item No.
<a href="#">ProCipitate™</a>	30 ml	P0050-30
<a href="#">ProCipitate™</a>	100 ml	P0050-100

### STORAGE

**ProCipitate™** is an aqueous suspension polyelectrolyte in distilled water. The reagent when not used must be kept sealed and stored at 4°C. **ProCipitate™** retains full activity when stored at 4°C for about 6 months. For long term storage, please contact us.

### Performance Characteristics

Protein	ProCipitate™: Sample	Removal
BSA, PBS @ 30 mg/ml	1 : 1	>99%
Human Serum	2 : 1	>90%
Nucleic Acid Recovery	ProCipitate™: Sample	Recovery
Calf Thymus DNA, $A_{260} = 1.00$	1 : 1	>95%
Total RNA, $A_{260} = 1.00$	1 : 1	>99%

## PROTOCOL

1. Resuspend **ProCipitate™** by shaking well prior to use.
2. Lyse sample to dissociate nucleic acids from histones and other proteins. Using wide bore or cut pipette tips, add the appropriate volume of **ProCipitate™** to deproteinize sample. Use Table above as a guide for volume addition or try several volume ratios starting with a maximum of 1 ml **ProCipitate™** to 1 ml of the sample (1 : 1 volume ratio).
3. Gently mix by inversion for 5 minutes at room temperature.
4. Centrifuge sample at 3000 x g for 15 minutes or microfuge at 16,000 x g for 5 minutes.
5. Recover purified nucleic acids contained in the supernatant.



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6. Continue with alcohol precipitation, DNA/RNA binding filter, or other suitable methods.  
 Note: Buffer condition may be at a moderately acidic pH and there may be a small volume dilution.

### **ProCipitate™** is Scaleable

The volumetric ratio of **ProCipitate** to sample can be adjusted up or down depending on the concentration of protein in the sample. Once established, these same ratios can be used to process volumes at any scale.

### Viral Nucleic Acid Isolation

Reference: Schwab, K.J., De Leon, R., and Sobsey, M.D., *Concentration And Purification Of Beef Extract Mock Eluates From Water Samples For The Detection Of Enteroviruses, Hepatitis A Virus, and Norwalk Virus by Reverse Transcription-PCR*, Applied and Env. Microbio, 61:531-537, 1995.

Another polyelectrolyte reagent, **Viraffinity™**, also can be utilized. Please contact us.

## References

### **ProCipitate™** appears in several articles and books on Food Safety

- ❖ **Foodborne Disease Handbook**, Second Edition, : Volume 2: Viruses: Parasites  
 By Y. H. Hui, Sayed A. Sattar, Wai-Kit Nip  
 "Viruses in the PEG eluants were precipitated...by an equal volume of **ProCipitate™**."
- ❖ **Health-related Water Microbiology**, Volume 27, Issues 3-4, Pergamon, 1993  
 "**ProCipitate™** was an effective method to purify the sample and dramatically improve virus detectability by RT-PCR."

### Patents

U.S. Patent Number 5,538,870, [Method for Preparing Nucleic Acids For Analysis And Kits Useful Therefore](#). This patent shows the beneficial effects of **ProCipitate™** in protocols which neutralize SDS with non-ionic detergents, are PCR compatible, and require no alcohol precipitation.

### Plasmids, Cosmids, BACs

Huang, G. M., et al, *A High-Throughput Plasmid DNA Preparation Method*, Analytical Biochem, 223:35-48, 1994.

Robert R. Klein, Daryl T. Morishige, Patricia E. Klein, Jianmin Dong; John E. Mullet. [High Throughput BAC DNA Isolation for Physical Map Construction of Sorghum](#). Plant Molecular Biology Reporter Dec 1998, Volume 16, Issue 4, pp 351-364



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Osoegawa, K. et al. *Bacterial Artificial Chromosome Libraries for Mouse Sequencing and Functional Analysis*. Genome Res. (2000) 10: 116-128.

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Quiniou SM; Katagiri T; Miller NW; Wilson M; Wolters WR; Waldbieser GC. *Construction and characterization of a BAC library from a gynogenetic channel catfish *Ictalurus punctatus**. Genetics, selection, evolution : GSE. 2003;35(6):673-83

Barbara J. Campbell, Jeffrey L. Stein, S. Craig Cary. *Evidence of Chemolithoautotrophy in the Bacterial Community Associated with *Alvinella pompejana*, a Hydrothermal Vent Polychaete*. Appl. Environ. Microbiol. Sept 2003; 69:9 5070-5078. doi:10.1128/AEM.69.9.5070-5078.2003

Chi, JX, et al. *Defining the orientation of the tandem fusions that occurred during the evolution of Indian muntjac chromosomes by BAC mapping*. Chromosoma. August 2005, Volume 114, Issue 3, pp 167-172

Stephanie M Cohen, Terrence S Furey, Norman A Doggett, David G Kaufman. *Genome-wide sequence and functional analysis of early replicating DNA in normal human fibroblasts*. BMC Genomics. 2006, 7:301 doi:10.1186/1471-2164-7-301

Amber E. Alsop, Andrew E. Teschendorff, Paul A.W. Edwards. *Distribution of breakpoints on chromosome 18 in breast, colorectal, and pancreatic carcinoma cell lines*. Cancer Genetics and Cytogenetics. Jan 2006. Vol 164, Issue 2 , Pages 97-109.

McDermott BM Jr; Asai Y; Baucom JM; Jani SD; Castellanos Y; Gomez G; McClintock JM; Starr CJ; Hudspeth AJ *Transgenic labeling of hair cells in the zebrafish *acusticolateralis* system*. Gene expression patterns. 2010;10(2-3):113-8

### Food Safety, Enteric Viruses and Environmental Sampling

Foodborne Disease Handbook, Second Edition, : Volume 2: Viruses: Parasites  
Y. H. Hui, Sayed A. Sattar, Wai-Kit Nip

By

Health-related Water Microbiology, Volume 27, Issues 3-4, Pergamon, 1993



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D'Souza, D. H. "Update on foodborne viruses: types, concentration and sampling methods." *Advances in Microbial Food Safety 2* (2014): 102.

### **Infectious Disease and Pathogen Detection, Paraffin-embedded Tissues**

Zoltan S., et al. [Detection of Mycobacterium avium Subspecies avium in Formalin-Fixed, Paraffin-Embedded Tissues of Captive Exotic Birds Using Polymerase Chain Reaction.](#) Journal of Zoo and Wildlife Medicine.1999;30:3:348-353

Miller, J.M., et al, [Polymerase chain reaction identification of Mycobacterium avium in formalin-fixed, paraffin-embedded animal tissues.](#) Journal of Vet.Diagnostic Invest.(1999)11:436-440

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### **Enzyme Removal (DNA)**

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### **Blood (DNA) PCR**

Krupey, J., et al, 100,000+ PCRs Possible from 10 ml Blood, poster Biotechniques Symposium, 2003.

### **Protein Enrichment**

J D Burton; R N Bamford; C Peters; A J Grant; G Kurys; C K Goldman; J Brennan; E Roessler; T A Waldmann. [A lymphokine, provisionally designated interleukin T and produced by a human adult T-cell leukemia line, stimulates T-cell proliferation and the induction of lymphokine-activated killer cells.](#) Proc of the Nat Acad of Sciences of the USA. (1994) 91(11): 4935-493.



## BIOTECH SUPPORT GROUP

### CONTACT US

**We welcome your questions and comments regarding our products.**

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