Water Quality Monitoring Project

In the

Barnegat Bay Watershed, WMA 13

New Jersey Department of Environmental Protection Water Monitoring and Standards

June, 2011 Revision 1 August, 2011 Revision 2 December, 2011

QUALITY ASSURANCE SAMPLING PLAN

Barnegat Bay Watershed, WMA 13

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1. **Project Name:** Barnegat Bay Sampling Project

2. **Requesting Agency:** NJDEP, Water Monitoring and Standards

3. Date of Project Requested: May, 2011

4. Date of Project Initiation: June, 2011

5. Project Officer: Jill Lipoti

6. Project Description:

Objective and Coverage

Current concerns regarding the use support within the Barnegat Bay Estuary derive in part from observed loss of sea grasses such as eel grass and widgeon grass, collectively referred to as Submerged Aquatic Vegetation (SAV), episodic occurrences of macro algae and brown tides, decline of hard clams, and increasing numbers of invasive species such as sea nettles. The full suite of stressors and biological/chemical/physical processes responsible for habitat alteration, loss of biological diversity and loss of support of designated uses are not entirely known. Alteration of the shoreline, hydrologic modification, resource harvesting, boating, the effects of the Oyster Creek nuclear generating facility and declining water quality are all suspected causes. With regard to water quality, available data indicate that there are areas within the Barnegat Bay where there are excursions from existing surface water quality standards, specifically with respect to dissolved oxygen and pathogen indicators. The Department has recently adopted narrative criteria for nutrients. The current study, as well as completed and other on-going studies, are expected to provide data that will assist in assessing the spatial extent of impairment with respect to these narrative nutrient criteria.

The purpose of this study is to provide water quality data to 1) determine the locations and extent of water quality impairments, 2) identify numeric criteria or loading targets for nutrients, including the need to revise existing SWQS to set the restoration endpoints (revisions may be needed to reflect a better link to supporting designated uses or where natural conditions would preclude support) and 3) calibrate and validate modeling tools that can be used to direct water quality restoration of the bay.

It must be noted that designing an effective data collection program depends on the objective, for example, the type of the impairment(s) to be addressed, the spatial extent and defining characteristics of the study area, the modeling approach (i.e., water quality, hydrologic, hydrodynamic), as well as specific models that must be supported. This sampling program has been prepared to maximize utility to address multiple objectives, while recognizing that there are unknowns at this juncture that may result in a need to revise the plan. In particular, it is not yet known which specific models will be used. This will require an adaptive approach as the project proceeds in order to obtain input from those who will be performing the modeling work.

In addition to characterizing pollutant loads, this study will further identify and characterize specific stressors of the Barnegat Bay ecosystem. Potential stressors may include the patterns of water circulation within the bay, shallow depth of water in the bay, thermal discharge from Oyster Creek, degree of biological productivity, sediment oxygen demand, turbidity and sediment toxicity.

The Department intends to work with Barnegat Bay partners to accomplish this intensive data collection effort. Critical to the success of this endeavor is the need to develop within this comprehensive Quality Assurance Project Plan measures to ensure that the laboratories conducting the analysis are certified for the work, that field staff (DEP and partners) are trained and that the laboratories are capable of handling the volume of samples. Therefore, the initial stage (first 6 months) of sampling effort is designed primarily to test out our capabilities, but will also provide data for condition assessment and assist with model development. This approach is necessitated in part because the monitoring can not be initiated in time to capture an entire growing season, the flow gauging stations will not be installed and operational until April, 2012 and monitoring needs may need to be adjusted once modeling tools are selected and the model needs are known. Another need to be met, which requires that this sampling program be initiated immediately, is to enhance baseline data to evaluate and capture the effect of implementing measures to reduce nutrient and other pollutant loads through stormwater infrastructure retrofits and fertilizer use restrictions. Once proof of concept has been achieved and adjustments made as needed, a minimum of a full year of monitoring will be required to develop modeling tools. If abnormal weather patterns prevail, it may be necessary to extend monitoring further in order to collect sufficient hydrodynamic, water chemistry, biological, flow, current, and sediment flux data for development of water quality/quantity models of the Barnegat Bay Estuary.

This field sampling study will address the following objectives:

- 1. Provide more comprehensive assessment of the relevant water quality conditions throughout the Barnegat Bay both spatially and temporally;
- 2. Provide water quality and biomass data to better quantify biological productivity and its impact on dissolved oxygen concentrations in the Bay.
- 3. Estimate the nutrient loadings into the bay and establish boundary conditions for the significant tributaries to the bay;
- 4. Provide nutrient concentration and loading data needed to evaluate the effects of nutrient load reduction scenarios.
- 5. Provide an understanding of the physical factors that may be affecting the bay water quality; such as the flushing rate, temperature, salinity and the depth of the bay. These factors play a major role in the physical, chemical and biological processes operating within the bay;
- 6. Collect sufficient data (minimum 24 months) to develop water quality and hydrodynamics models;
- 7. Collect data that will capture seasonal variability as well as variability between years;
- 8. Collect sediment toxicity data to assess if biological changes loss of SAV, decline in hard clams, and increase in invasive species are caused by sediment toxicity.

This proposed monitoring program will consist of the following components:

- 1. Continuous in-situ water quality monitoring,
- 2. Grab water quality sampling,
- 3. Two 5-day intensive sampling events;
- 4. Sediment characterization and toxicity monitoring;

¹ Partners are approved to conduct field analysis and sample collection only for Barnegat Bay project work until June 20, 2012

- 5. Flow monitoring;
- 6. Bathymetric survey

Phase 1 will provide a proof of concept for the monitoring plan, including the partnership approach. Monitoring partners will be asked to sign an MOU in which they agree to follow the procedures set forth in this OAPP. Sampling will be conducted under various conditions and frequencies. Phase 2 will continue the program for a minimum combined duration of 24 months. However, the sampling period may need to be extended if the conditions experienced are deemed not representative. This Quality Assurance Sampling Plan is developed for Phase 1 of this sampling plan, an addendum will be prepared to reflect refinements for Phase 2 and constitute the Phase 2 Quality Assurance Sampling Plan.

7. Sampling Network and Design Rationale

7.1-**Continuous in-situ water quality monitoring:**

Physical and biological parameters such as temperature, dissolved oxygen (DO), DO % saturation, pH, conductivity (salinity), and turbidity in the Barnegat Bay are required for water quality model simulation and, to some extent, for the hydrodynamic model (especially temperature and salinity). Continuous monitoring multi-parameter probes will be deployed at the proposed monitoring sites in Barnegat Bay. Table 1 specifies the list of parameters, frequency of collection and the number of sites for each phase of sampling. Table 2 identifies the bay sampling locations and Figure 1 shows the location of all of the proposed sampling sites.

Table 1: Continuous Monitoring Plan: Monitoring Parameters and Frequency of Collection

	Continuous Monitoring Plan* Bay									
Sampling Stations	Sampling Type	Sampling Matrix	Parameters	Frequency						
Phase 1 sampling 4 sites, as identified in Table 2 and Figure 1 below	Continuous monitoring probes at mid depth ² using 4-6 buoy-located devices and 1 device housed within a fixed station	Aqueous	Dissolved Oxygen concentration (DO), pH, Temperature, Turbidity, Conductivity, Salinity ³ , Chlorophyll-a; (possible deployment of NO ₃ probe at Mantoloking using a buoy provided by	Measurements every 15 minutes. Duration of sampling is expected to be continuous for a minimum of 24 months.						

² Sampling sites where depth is greater than 12 feet, two samples will be taken at 1/3 and 2/3 depth.

³ Salinity to be calculated from conductivity

Phase 2 sampling 6 buoy locations and 1 fixed station, as identified in Table 2 and Figure 1 below	sampling		USGS)	
		Tributary		
Sampling Stations	Sampling Type	Sampling Matrix	Parameters	Frequency
Toms River near Thomas River (USGS 01408500)	probes located within the existing gauging station	Aqueous	Dissolved Oxygen concentration (DO), pH, Temperature, Turbidity, Conductivity, Nitrate/Nitrite	Measurements every 15 minutes. Duration of sampling is expected to be continuous for a minimum of 24 months.

^{*} A separate QAPP has been developed for the Department's continuous monitoring efforts

Table 2: List of monitoring sites within the Barnegat Bay

Project ID	Site Description	Type: Grab/ Buoy/Fixed	Longitude	Latitude	Site Reference ID	Site Partner
BB01	Barnegat Bay at Mantoloking	G,F	-74.054320	40.038320	USGS- 01408168	Monmouth University
BB02	Barnegat Bay between Silver Bay and Goose Creek	G,B	-74.098470	39.9776200	BMWM1622	Monmouth University
BB03	Barnegat Bay by Route 37 Bridge	G	-74.101530	39.9481700	BMWM1629B	Monmouth University
BB04	Barnegat Bay near the Mouth of Toms River	G	-74.110140	39.9376200	BMWM1623B	Monmouth University
BB05	Barnegat Bay above Cedar Creek	G,B*	-74.112910	39.8845600	BMWM1645E	EPA
BB06	Barnegat Bay below Cedar Creek and above Forked River	G	-74.102080	39.8526200	BMWM1651D	EPA
BB00	Barnegat Bay below Oyster Creek and above Barnegat	G,B	71.102000	33.0320200	BITWITTOSTE	EPA
BB07	Inlet		-74.153190	39.7926200	BMWM1691A	
BB08	Barnegat Bay by Barnegat Inlet	G	-74.108014	39.7633528	MU-Barnegat Inlet	EPA.
	Barnegat Bay below Barnegat Inlet and close to	G,B*				DEP- Leeds Pt.
BB09	Long Beach		-74.147920	39.7426200	BMWM1674B	
BB10	Barnegat Bay by Route 72 Bridge	G	-74.206530	39.6609500	BMWM1703C	DEP- Leeds Pt.
BB11	Barnegat Bay above Westecunk Creek	G	-74.235700	39.6254000	BMWM1719E	DEP- Leeds Pt.
BB12	Barnegat Bay in Little Egg Harbor	G,B	-74.268750	39.5815100	BMWM1834A	DEP- Leeds Pt.
BB13	Barnegat Bay near Tuckerton Creek	G	-74.324590	39.5690100	BMWM1818D	DEP- Leeds Pt.
BB14	Little Egg Harbor Inlet near Beach Haven Heights	G,B	-74.297370	39.5112300	BMWM1824B	DEP- Leeds Pt.

* Monmouth University will use buoy to collect real time data at this station.

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In addition to this project, Monmouth University will be collecting data using buoys at 4 locations within Barnegat Bay, addressed in a separate QAPP (Quality Assurance Project Plan for Water Quality Monitoring in Coastal Ocean and Monmouth Counties Using Data Loggers Monmouth University School of Science Urban Coast Institute). The data collected in the Monmouth University study will be used to supplement the study described herein.

7.2- Field Parameters

Field parameters, pH, water temperature, dissolved oxygen and specific conductance, will be measured on site. Collected turbidity samples will be measured at a project field station by Bureau of Freshwater and Biological Monitoring staff certified for turbidity measurements. Sample filtration for dissolved parameters will also be conducted by Bureau of Freshwater and Biological Monitoring staff and Bureau of Marine Water Monitoring staff. At each sampling location, analyze immediately parameters (i.e. pH, specific conductance, salinity (calculated from specific conductance), dissolved oxygen, and turbidity) will be taken using handheld meters or multi parameter sensors.

7.3- Grab Water Quality Sampling:

Grab samples will be collected at the Barnegat Bay locations listed on Table 2 and additionally at tributary sites listed on Table 5. Locations are shown in Figure 1. The proposed frequency varies by phase and season and is listed in Table 3. The sampling period may be extended in response to information obtained during the initial phase of sampling or if conditions experienced are not considered to be representative. Samples will be collected in accordance with approved field sampling procedures and analyzed in certified laboratories. As indicated, the water quality parameters to be sampled will be driven, in part, by the modeling tools selected, but are expected to include those listed in Table 3.

All water quality grab samples will be collected following procedures found in "NJDEP Field Sampling Procedures Manual, August 2005". Sampling locations have been marked and verified with GPS. In addition NJDEP staff and project partners will utilize detailed site sketches to locate the sampling location on the first and subsequent visits. The freshwater tributary locations samples will be collected as center of flow grab samples. At tributary locations greater than 20 ft wide specific conductance measurements were made along a transect and it was determined that at all locations the stream is well mixed and that a center of flow grab sample would be representative of the water quality at that location. Because none of the tributary monitoring locations is greater than 12 ft deep, samples will be collected at a depth of 1 ft. The project began with in bay water quality samples taken as surface grab samples only, at the designated locations. In order to determine if the bay was well mixed for TSS and turbidity the collection of a bottom grab sample for these parameters was added. Therefore starting with the 5th (8/25/2011) sampling event a surface and bottom grab sample will be taken at the bay sampling locations. Bottom grab samples will be taken using a Kemmerer sampler approximately 6in-1ft above the bottom sediment. Surface grab samples will be taken first followed by the bottom grab sample. All samples will be taken to a field location for filtration and transportation as reflected in Table 4 and Figures 2-4. All sample containers are being supplied by the DEP and only these sample containers can be used for the project. In addition, for field testing for pH, the DEP will be supplying a single lot of certified primary pH buffers for use in the calibration of the meters. All sample containers must be transported on ice in coolers to preserve the integrity of the samples and maintain sample temperature at greater then freezing and less then 6°C. Necessary acids as preservative will be added at the field station (FREC) or Leeds Point Lab.

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Table 3: Grab Samples: Monitoring Parameters and Frequency of Collection in Barnegat Bay and its Tributaries

		Monitoring	Parameters and Frequen	ncy of Collection		
Sampling Stations	Sampling Type	Sampling Matrix	Parameters	Lab		uency
			Total Suspended Solids (TSS) ³	Leeds Point	Phase one:	Phase two:
			Chlorophyll-a (w/species ID)* Total Nitrogen (TN) Dissolved Total N	Leeds Point Leeds Point Leeds Point		
			Dissolved Ammonia	Leeds Point		
			Dissolved Nitrate+Nitrite	Leeds Point	_	
			Total Phosphorus (TP)	Leeds Point		one sample
			Dissolved Total Phosphorus	Leeds Point		twice a month
			Dissolved Ortho-P	Leeds Point		January
			Total Organic Carbon	EPA Edison and Maryland/Leeds ¹	one sample	2012 through
			Dissolved Organic Carbon	EPA Edison and Maryland/Leeds ¹	every two weeks	March 2012, one
Locations	Surface		Dissolved Inorganic Carbon ²	Maryland ²	from June 2011	sample per week from
are	grabs and		Alkalinity	EPA Edison	through	April
identified in Tables 2 and	bottom grabs* ³	Aqueous	Unfiltered Total BOD5	OCUA	September 2011 and	1,2012 through
4.	graus		Unfiltered CBOD5 (nitrification inhibited and no seed or pH		then once	September
			adjustment)	OCUA	per month from	30, 2012 and
			Unfiltered CBOD20 (nitrification inhibited and no seed or pH		October	resuming
			adjustment)	OCUA	2011	one sample
			Total Si	EPA Edison	through December	every two weeks in
			Dissolved Si	EPA Edison	2011	October
			Turbidity ³	Leeds/BFBM		2012 through
			Secchi depth*	Field		December
			Transmissometry*	Field		2012
			pH ³	Field	_	
			Dissolved Oxygen (DO) ³	Field	_	
			DO Saturation ³	Field	_	
			Temperature ³	Field	_	
			Conductivity (Salinity ³)	Field		

^{*}omit in tributaries

Note: 1. EPA lab will perform the Freshwater sample analysis and Maryland/Leeds will perform the salt water sample analysis. Where Maryland/Leeds is the denoted lab, Maryland will be used until the appropriate equipment is on-line at Leeds. There will be duplicative sampling for three events to allow a determination of the degree of variability between the two labs and two methods of analysis. Once Leeds has the equipment, the method that will be followed is SM 5310C.

^{2.} Only for salt water sample. DIC for the freshwater sample will not be analyzed but will be calculated based on the Alkalinity results for the freshwater samples.

3 For bay sampling locations a bottom grab sample was collected from sampling event 8/25/11 and analysis for limited parameters, beginning in January 2011 and forward, this sample will be analyzed for TSS, Chlorophyll A, Turbidity, TP, TN, TOC, and Total Silica . In-situ measurements of pH, dissolved oxygen (DO),, DO saturation, temperature and conductivity will also be conducted at depth where the bottom grab samples are collected.

Table 4. THE ANALYTICAL METHOD TABLE

	1able 4. THE ANA	LIII	AL N	IETHOD	TABLE			11-1-1:	
Org	Parameter		Prep	Code	Method	Container	Preservative	Holding Time	Bottle
NJDEP BFBM		- 0			0110100 B			40.1	- .
_	Turbidity Total Suspended	FwSw	U	Turb	SM 2130 B USGS I-3750-	50 mL centrifuge tube	Ice, 4°C	48 hours	T1
	Solids	FwSw	U	TSS	85	Amber 500 mL HDPE	Ice, 4ºC	24 hours	L1
	Chlorophyll a (bay only)	Sw	U	Chla	SM 10200-H	Amber 500 mL HDPE	Ice, 4°C	24 hours	L1
	Total Nitrogen	FwSw	U	TN	USGS I-4650- 03	50 mL centrifuge tubes	Ice, 4°C	28 days	L2
	Total Phosphours	FwSw	U	TP	EPA 353.4	50 mL centrifuge tubes	Ice, 4°C	28 days	L2
	Dissolved Ammonia	FwSw	F	DNH3	SM 4500- NH3(G)	50 mL centrifuge tubes	2 ml 3.5% Phenol	14 days	L3
q	Dissolved Nitrite + Nitrate	FwSw	F	DNO3	EPA 353.4	50 mL centrifuge tubes	Ice, 4°C	28 days	L4
NJDEP Leeds Point Lab	Dissolved Orthophosphate	FwSw	F	DPO4	EPA 365.5	50 mL centrifuge tubes	Ice, 4°C	28 days	L4
» Po	Dissolved Nitrogen	FwSw	F	DN	USGS I-4650- 03	50 mL centrifuge tubes	Ice, 4ºC	28 days	L4
Leeds	Dissolved Phosphours	FwSw	F	DP	EPA 353.4	50 mL centrifuge tubes	Ice, 4°C	28 days	L4
굡	Total Organic Nitrogen	FwSw	NA	TON	Calculated	NA	NA	NA	NA
NJD	Dissolved Organic Nitrogen	FwSw	NA	DON	Calculated	NA	NA	NA	NA
	Dissolved Organic Phosphorus	FwSw	NA	DOP	Calculated	NA	NA	NA	NA
	Particulate Organic Nitrogen	FwSw	NA	PON	Calculated	NA	NA	NA	NA
	Particulate Phosphorus	FwSw	NA	PP	Calculated	NA	NA NA	NA	NA
	Dissolved Organic Carbon	Sw	F	DOC	SM 5310 C	Glass, 250ml	Conc. H2SO4, pH<2	28 days	L7
	Dissolved Inorganic Carbon	Sw	F	DIC	SM 5310 C	Glass, 250ml	Conc. H2SO4, pH<3	29 days	L7
							Conc. H2SO4,		
	Total Organic Carbon	Sw	U	TOC	SM 5310 C	Glass, 250ml	pH<3	28 days	L8
	Biochemical Oxygen Demand (5 day)	FwSw	U	BOD5	SM 5210B	2 L HDPE	Ice, 4°C	48 hours	O1
OCUA	Carbonaceous Oxygen Demand (20 day)	FwSw	U	CBOD5	SM 5210B	2 L HDPE	Ice, 4ºC	48 hours	02
	Carbonaceous Oxygen Demand (5 day)	FwSw	U	CBOD20	SM 5210B	2 L HDPE	lce, 4ºC	48 hours	O3
7	Alkalinity	FwSw	U	Alk	SM 2320 B	250 mL HDPE	Ice, 4°C	14 days	E1
ion	Total Silica	FwSw	U	Si	EPA 200.7 rev 4	125 mL HDPE	Conc. HNO3, pH<2	6 months	E2
eg n L	Dissolved Silica	FwSw	F	DSi	EPA 200.7 rev 4	250 mL HDPE	Conc. HNO3, pH<3	7 months	E3
USEPA Region 2 Edison Lab	Total Organic Carbon	Fw	U	TOC	SM 5310 B	250 mL HDPE	Conc. H2SO4, pH<2	28 days	E4
ISN B	Dissolved Organic Carbon	Fw	F	DOC	SM 5310 B	250 mL HDPE	Conc. H2SO4, pH<3	29 days	E5

a: Carbon analysis in saltwater samples was performed by Maryland University Laboratory for Phase 1.

Table 5: Tributary Sampling Locations

		Description			Flow-	Flow	Water Quality
					Measurement	Site	Site
Station ID	Site #		LATITUDE	LONGITUDE	Type	Partner	Partner
		North Branch			Extrapolate from	NA	Brick
		Metedeconk R near			existing gage		MUA
USGS-01408123	BT01	Laurelton	40.081648	-74.151811			
					Gage (new)		Brick
		SB Metedeconk			Measure phase 1		MUA
		River near			(if wadeable and		
		Laurelton			until relationship		
		(Chambers Bridge			established with		
USGS-01408152	BT02	Rd)	40.078763	-74.156729	gage site)	USGS	
		Toms River near			Gage (existing)	NA	DEP
USGS-01408505	BT03	Toms River	39.976389	-74.218333			
		Wrangle Brook near			Measure	NJDEP	BBP
USGS-01408640	BT04 ¹	South Toms River	39.952854	-74.218515			
		Jakes Branch at			Measure	NJDEP	OCHD
		South Toms River					
USGS-01408710	BT05	(route 619)	39.935418	-74.211554			
		Cedar Creek at			Gage (existing)	NA	OCHD
USGS-01409000	BT06	Lanoka Harbor	39.868887	-74.17043			
		NB Forked R at			Measure	NJDEP	MATES
USGS-01409055	BT07	Forked River	39.836035	-74.196013			
		Middle Br Forked			Measure	NJDEP	MATES
		River (upstream Rt		_,_,,			
BFBM000165	BT08	9)	39.828238	-74.201971	2.5		
		South Br Forked			Measure	NJDEP	DEP
DEDM000166	DTOO	River (upstream Rt	20.02020	74 202120			
BFBM000166	BT09	9 @ JCPL)	39.82038	-74.203128	Gage (new)		DEP
		Oyster Creek (upstream Rt 9 @			C \ /		DEP
BFBM000167	BT10	JCPL)	39.810584	-74.204626	Measure Phase 1	NJDEP	
DFDWI000107	D110	Mill Ck at	39.810384	-74.204020	Gage (new)	NJDEP	
		Manahawkin (Bay			• ,		Pinelands
USGS-01409210	BT11	Avenue)	39.695405	-74.259527	Measure Phase 1	USGS	rinelands
0303-01407210	DIII	Westecunk Ck at	37.073403	-14.237321	Measure	0303	Pinelands
		Railroad Ave at			ivicasuit	USGS	1 illelands
USGS-01409281	BT12	West Ck	39.640297	-74.30797		0000	
0505-01-07201	D112	Tuckerton Creek at	37.070291	-17.30171	Measure		DEP-
USGS-01409310	BT13	Tuckerton*	39.60274	-74.34163	Measure	USGS	Leeds Pt.
		2011 than suspended d				0000	Lecus I t.

^{*} From June through August 2011, then suspended due to reconstruction of dam

Note: 1. For the initial sampling event 6/6/2011 BT04 was sampled at the incorrect location (Lat 39.960999, Long-74.228078), the limited data available for this station appears under the Site ID BT04a

In addition, NJDEP will deploy a temperature data logger in the cooling water channel at the Oyster Creek Generating Station. Measurements will be made every 0.5 hrs to document general daily fluctuations and brief, but potentially significant, abrupt changes in water temperature. The data logger will be deployed during the summer of 2011. A separate QAPP developed for ambient water temperature monitoring will apply to this activity.

Table 6: Dates and time frame of sampling collection

Phase 1 Sampling Event	<u>Dates</u>	Planned Bad Weather Date	Date Sampling Was Performed	Sample Collection Times Trib	Sample Collection Time Bay
1	6/6/2011	NA	6/6/2011	8:00-12:00	8:00-12:00
2	6/23/2011	NA	6/23/2011	8:00-12:00	8:00-12:00
3	7/5/2011	NA	7/5/2011	8:00-12:00	8:00-12:00
4	7/21/2011	NA	7/21/2011	8:00-12:00	8:00-12:00
5	8/8/2011	NA	8/8/2011	8:00-12:00	8:00-12:00
6	8/25/2011	NA	8/25/2011	8:00-12:00	8:00-12:00
7	9/15/2011	NA	9/15/2011	8:00-12:01	8:00-12:01
8	9/26/2011	NA	9/26/2011	8:00-12:02	8:00-12:02
9	10/13/2011	NA	10/13/2011	8:00-12:03	8:00-12:03
10	10/24/2011	NA	10/24/2011	8:00-12:04	8:00-12:04
11	11/10/2011	NA	11/14/2011	8:00-12:05	8:00-12:05
12	12/8/2011 *	NA	12/12/2011	8:00-12:06	8:00-12:06
Phase 1 Extended	Dates	Planned Bad Weather Date	<u>Date Sampling</u> Was Performed	Sample Collection Times Trib	Sample Collection Time Bay
1	1/10/2012	1/17/2012		8:00-12:00	8:00-12:00
2	1/26/2012	1/31/2012		8:00-12:00	8:00-12:00
3	2/7/2012	2/14/2012		8:00-12:00	8:00-12:00
4	2/23/2012	2/29/2012		8:00-12:00	8:00-12:00
5	3/6/2012	3/13/2012		8:00-12:00	8:00-12:00
6	3/22/2012	3/29/2012		8:00-12:00	8:00-12:00

Dates where changed due to scheduling conflicts and severe inclement weather

7.4- Two 5-Day intensive sampling events: (PLACEHOLDER)

In addition to the periodic grab sample program identified in Task 2, two 4-day intensive surveys are also planned during the phase 2 sampling program. Grab samples will be collected during each of the two 4-day events, at a frequency of six (6) per day. These intensive sampling events are needed to properly calibrate the water quality model. The sampling parameters and sites will be described in an addendum to this QAPP. The first sampling event should be accomplished within July of 2012 and the second event within August 2012. The intensive sampling task may be adjusted after review of the Phase 1 data.

7.5- Sediment Monitoring: (Placeholder)

Sediment can also impact the dissolved oxygen demand in the Bay through biological and chemical activities. Sediment toxicity may be a cause of changes observed in SAV, hard clam populations and invasive species. To characterize the effects of sediment relevant to water quality model development and simulations, physical and chemical makeup of the sediment must be determined see Table 7.

Table 7 - Sediment physical and chemical characterization monitoring:

Monitoring Parameters and Frequency of Collection							
Sampling Sites	Sampling Type	Parameters	Frequency				
At a minimum, 3 sites within the Bay and at mouth of Toms River	Core and Grab—	 ✓ SOD ✓ Dissolved Inorganic P, N, C ✓ Particulate Inorganic P, N, C ✓ Particulate Organic P, N, C ✓ Dissolved Organic C ✓ Grain size and type ✓ Flux - Sulfide, NO3, NH3 and DIP ✓ Pore water NH3, NO3, DIP, H₂S, Si, ✓ TN ✓ TP ✓ Sulfide 	Monthly samples - July through September				

<u>Placeholder: Wave & Re-suspension Monitoring</u>: (Phase 2, depends on model needs)

Measure shallow water wave energy and sediment re-suspension at key points. Move an instrumentation package weekly to different shoals/beaches to obtain distribution of data. Consider adding UGEMS sediment erodibility measurements on cores. Allow for calibration of a sediment re-suspension of modeling component. Model selection will determine need for this component.

Placeholder: Side Scan SONAR& Sediment Flux

7.6- Flow monitoring:

In addition to existing gages, 9 additional gages will be installed to measure flow. These data are necessary inputs for the hydrodynamic and water quality models. The locations of existing and new gages are presented in Table 8 and shown in Figure 1. At tributary locations where gages are absent, flow will be measured using hand held equipment such as SONTEK Flow Tracker (or equivalent). Flow measurement SOP is available in Flow Tracker Handheld ADV Users Manual (SonTek/YSI 2009 FlowTracker Handheld ADV User's Manual Firmware Version 3.7). Discharge measurements at higher stages that can not be waded at the SB Metedeconk River at Chambers Bridge Road near Laurelton will be made from the bridge or a pulley system. A Teledyne RDI StreamPro Acoustic Doppler Current Profiler (ADCP) mounted to a tethered boat will be pulled across the stream from the downstream side of a bridge or on a pulley system from bank to bank (Gotvald and Oberg, 2009). A minimum of four transects are made across the channel. The ADCP transmits acoustic pings that record the velocity and depth of water. A Bluetooth wireless link from the ADCP transmits depth, distance and velocity data every second to a field computer. The field computer calculates the discharge for each transect. The discharge from the 4 transects are averaged. The USGS quality assurance plan for discharge measurements using ADCPs is published in

Oberg and others, 2005. References are available online at http://pubs.er.usgs.gov/usgspubs/sir/sir20055183. and http://pubs.er.usgs.gov/usgspubs/fs/fs20083096

Table 8: Existing and New Gauging Stations

Station Description	Latitude	Longitude	Type
Westecunk Creek at Stafford Forge NJ	39.666667	-74.320278	Existing-Trib
Cedar Creek at Lanoka Harbor NJ	39.8675	-74.169167	Existing-Trib
North Branch Metedeconk River near Lakewood			
NJ	40.091667	-74.1525	Existing-Trib
			New-
Point Pleasant Canal at Point Pleasant, NJ	40.070278	-74.059722	outlet/inlet
Barnegat Bay at Mantoloking Bridge at			
Mantoloking	40.04	-74.057222	New - in bay
Barnegat Bay at Route 37 Bridge near Bay Shore,	39.946111	-74.103056	New - in bay
			New-
Barnegat Inlet at Barnegat Light, NJ	39.766389	-74.099167	outlet/inlet
Barnegat Bay at Route 72 Bridge near Ship			
Bottom	39.663333	-74.206944	New - in bay
			New-
Little Egg Harbor Inlet near Beach Haven Heights	39.5075	-74.3075	outlet/inlet
Oyster Creek near Brookville, NJ	39.798333	-74.250556	New-Trib
S.B. Metedeconk River near Lakewood, NJ	40.085833	-74.185556	New-Trib
Mill Creek at Manahawkin, NJ	39.695278	-74.26	New-Trib

7.7- Bathymetric Survey (Placeholder)

The existing bathymetry was prepared in 1934-36 and predates significant events, such as modifications to the Barnegat Bay inlet. Limited recent bathymetric data is available in the vicinity of Barnegat Inlet. These data show that while some locations have experienced minor change, others have undergone significant change. Verification of the bathymetric configuration is needed to ensure reliable modeling results. It is also necessary to characterize sediment for sediment transport modeling. High resolution bathymetry will enhance the accuracy of the circulation model, which in turn, will enhance the ability to simulate sediment transport and other important parameters, e.g., salinity and temperature. A sampling project for measurement of Barnegat Bay bathymetry, bottom-sediment type and shallow structures is proposed using side-scan sonar swath bathymetry, sub bottom profiler, and sediment sampling. This method will be applied in deeper waters. Spot verification using equipment yet to be specified will be applied in shallow waters where the sonar equipment cannot be used.

Data Usage:

Water quality data sampled under this project will be used to identify assess water quality impairment and populate, calibrate and validate modeling tools along all other quality assured historic data. All sampling procedures must be in conformance with NJDEP or USGS (URL http://water.usgs.gov/owq/FieldManual/index.html) field sampling procedures as well as other applicable guidance. If a method or procedure requires change and is not contained in Table 3 and Table 4, this information should be brought to the attention of the signatories of this QAPP through writing and needs approval prior to being used.

8. Reports:

All water quality data collected, locations of final sampling sites, and related field notes should be entered in the New Jersey Water Quality Data Exchange (WQDE). Data quality assurance will occur at NJDEP Bureau of Freshwater and Biological Monitoring and Bureau of Environmental Analysis and Restoration using protocols found in USGS open file Report 02-383 "Methods for Quality Assurance Review of Water Quality Data in New Jersey".

9. Project area:

Watershed project area covered under this project is the Barnegat Bay Watershed in WMA 13 (see Figure 1 for the spatial extent of the study). The GIS map provided identifies proposed monitoring locations, dischargers, and approximate head of tide.

10. Data Representativeness

The same methods and techniques will be used by all field collection staff. Office of Quality Assurance will be on-site for the 1st sampling event to review and ensure that all samples are collected per the QAPP. Any deviations from the QAPP will be documented and will be resolved prior to the next sampling event

11. Data Validation:

Method blank (lab), equipment blank, duplicate, and replicate samples will add approximately 10 percent more to the total number of samples collected. A midpoint project review is scheduled for Sept 2011. At that time all aspects of the sample collector procedures will be evaluated and revisions incorporated, as needed. An addendum to this plan will be issued at that time detailing any changes. The data is verified using the replicate data percent difference discussed above. The data is validated using the QC data. The QC sample should fall between two standard deviations at the 95th percentile confidence level to be valid. All laboratory and field spikes should be with between 80-120 %. Water quality results will be assessed against available, historical water quality data from the locations monitored. Data will also be assessed using USGS Open-File Report 02-383 "Methods for Quality Assurance Review of Water-Quality Data in New Jersey ". That report provides information on standard ranges of specific parameters in New Jersey streams and standard relationships between specific parameters. All data collected will be provided to NJDEP and WM&S staff will perform the data validation process. Data that can not be confirmed by these reviews or explained by circumstances (i.e. heavy rain, drought) or project QA data will be classified as questionable by NJDEP. In addition, quality assurance protocols will be used by EPA Edison, OCUA, Leeds Point, the Maryland Lab and BFBM for the data validations under the supervision of a quality assurance officer.

12. Data Quality Requirements

Continuous Data Quality

Data recorders are calibrated and programmed within 24 hrs of each deployment following the manufacturer's manual. Duplicate DO measurements are made at the time of meter deployment and meter retrieval with a second meter, calibrated on site. Comparative DO readings not within the stated accuracy of the meters used will be reviewed against historical water quality data from that site as an additional quality review step. Data outside the stated accuracy of the meters used in the comparative readings and outside the historical range for DO at that location will not be used. At each sampling event water quality grab samples will be taken at the location of the continuous meters and analyzed for the parameters listed in Table 4. This data will be utilized to validate the data collected by the continuous meters.

Field Quality Assurance and Quality Control

NJDEP and Partner groups field staff will be approved by DEP's Office of Quality Assurance for field measurements, which include: specific conductance (Wheatstone Bridge, SM 2510 B), dissolved oxygen (electronic SM 4500-OG), pH Electronic SM 4500-HB) and temperature (Thermometric). Project staff will follow manufacture's manuals regarding calibration and operating procedures for specific meters. Results of daily pH calibrations, D.O. air calibrations and specific conductance calibrations will be recorded on field calibration forms. Weekly temperature ASTM- QC checks and Winkler D.O. checks are also recorded. Turbidity samples will be analyzed at one of the laboratories by NJDEP field staff who are certified for the measurement of turbidity (Nephelometric, SM 2530 B), The marine sample field quality control will consist of analyzing in the laboratory, the remaining sample not used for filtration for salinity; in addition, a dissolved oxygen Winkler titration sample will be collected. This data will be used to validate the data collected by sensors in the field. The Winkler titration sample must be protected from the intrusion of atmospheric oxygen and needs to be analyzed prior to the validation for the salinity.

The Laboratories participating in this project are state or National Environmental Laboratory Accreditation Program (NELAP) certified to perform the parameters conducted for ambient water quality monitoring and will follow the Laboratory methods as outlined in Appendix A. Any changes to the methods used must be preapproved by the DEP before sample testing continues.

The field meters or multi parameter sensors will be calibrated using manufacturer specifications. Calibration and verification will be performed with the following:

Temperature

Temperature thermistors are factory calibrated. Thermistors must be checked against a National Institute of Standards and Technology (NIST)-certified thermometer on a quarterly basis. If not found to be accurate within + 0.5 °C of the certified thermometer an offset value will be applied to correct the reading or if drift is continuing to take place sensor will be replaced. Any change will be noted in the calibration log. Temperature units will be degrees Celsius (°C). On August 17, 2011, the temperature monitoring devices were calibrated by DEP personnel against a NIST certified thermometer. This calibration must be repeated on August 17, 2011 and quarterly thereafter through the duration of the project.

Duplicate testing is required once every 20 samples tested.

Salinity/Specific Conductance

Specific conductance is calibrated using a factory prepared conductivity standard with a value of 50 mS/cm or a 35 ppt salinity standard for the marine samples and 1.412 mS/cm for the fresh water locations. Specific conductance units will be mS/cm, Salinity will be expressed in parts per thousand (ppt). For sensor verification, another standard from a different source will be analyzed. The calibration must be checked in the measure mode with a standard. The required accuracy is that the calibration check data must be within 1% of the true value of the standard used to be acceptable for analysis. Duplicate testing is required once every 20 samples tested.

Dissolved Oxygen

Calibration of a DO meter at 100 percent oxygen saturation is made by adjusting the meter reading for air saturated with water vapor, as per the manufacturer's instructions. Sensors will record both Dissolved Oxygen (DO) milligrams/liter (mg/l) and DO percent saturation (%). Samples for the Winkler titration will be collected at the marine water sites for sensor verification. Each week of use the DO meter must be verified against a Winkler titration procedure. The accuracy required between the reading from the DO meter and the results of the Winkler test must be within +/- 0.3 mg/L of each other to be acceptable.

Duplicate testing is required once every 20 samples tested.

pН

Most multi-meters require the use of a three point calibration with 4, 7 and 10 pH buffers. A three point calibration is the preferred approach to a quality calibration. All calibrations must be performed with the buffer solutions supplied by the DEP for consistency and must meet the accuracy requirement of being within 0.05 s.u. of the true value of the buffer used to be considered acceptable. A two point calibration can be performed using 7 and 10 buffers for the marine locations and 4 and 7 buffers for the fresh water locations as long as a calibration check (with the instrument in the measure mode) is conducted with the second of the two buffers used for calibration (i.e. 10 buffer for the marine and 7 buffer for the freshwater). The required calibration check result must be within 0.10 s.u. of the true value of the buffer used. Every three hours of use the meter must be checked with the calibration check buffer and must be accurate to 0.2 s.u. of the true value to be considered acceptable for continued use. The field staff may also recalibrate the meter at each site as an alternative to the three hour calibration check requirement.

Duplicate testing is required once every 20 samples tested.

Millivolt readings are also taken as a check of probe performance. For sensor verification, another certified pH buffer from a different source will be analyzed.

Turbidity

Turbidity samples will be analyzed at one of the field laboratories (FREC or Leeds Point). Calibration of a turbidity meter will be accomplished by a 2 point method using an YSI produced microbead synthetic turbidity standard (123NTU) and deionized water (0NTU) each day of use. Turbidity units will be (NTU).

Duplicate testing is required once every 20 samples tested.

A formazin standard or a standard from a different source will be analyzed for sensor verification.

Filtration Quality Control

Filtration quality control will consist of analyzing a filtration blank, that will be deionized water run through the pump tubes and the filter, preserved and analyzed as the other nutrient samples, prior to the filtration of a sample. Between each sample, the pump tubes will be flushed with a cycle of deionized water/ 10% HCl/ deionized water. The filtration blank will be repeated after roughly half (13 samples) and at the end of the processing of 27 samples. This will ensure the validity of the data and the pump cleansing process. In addition, 2 filtration spikes and replicates will be performed for each sample run. The spike will consist of adding a known amount of analyte to a volume of sample, and the sample will be filtered and processed as the other samples, the spike will ensure that there are no interferences, loss of analyte or contamination of the sample. Filtrations must be performed within 8 hours of sample collection or sooner and the time of filtration will be documented in the laboratory records or on the chain of custody form for the project.

Table 9

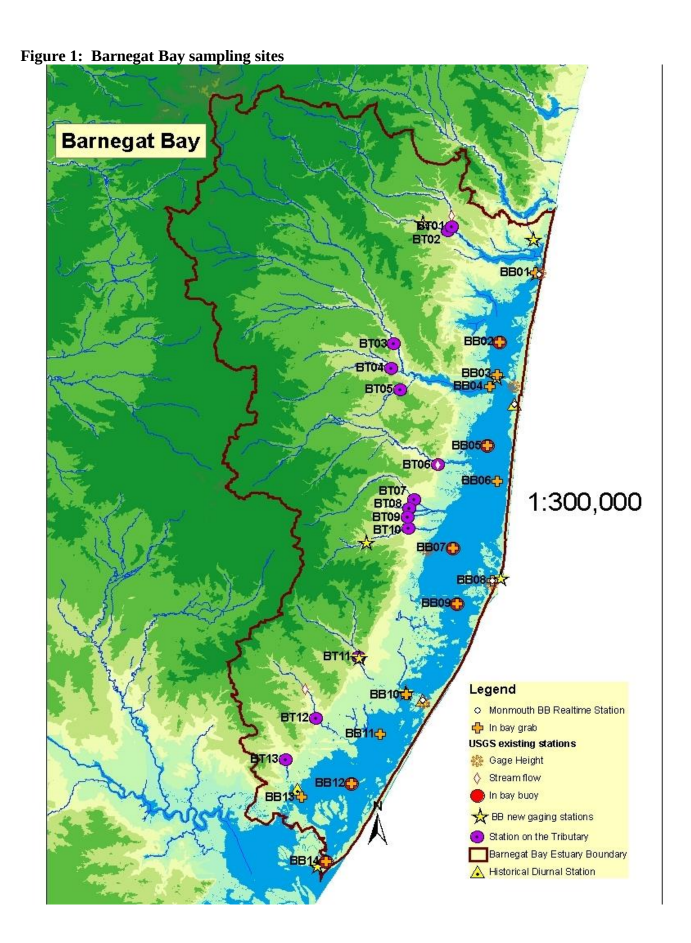
<u>Table 9</u>					Lab		
					Reporting		Holding
Lab	Parameter		Prep	Code	Limit	Method	Time
FREC/Leeds Point	Turbidity	FwSw	U	Turb	0.1 NTU	SM 2130 B	48 hours
						USGS I-3750-	
Leeds Point	Total Suspended Solids	FwSw	U	TSS	1.0 mg/l	85	24 hours
Leeds Point	Chlorophyll a (bay only)	Sw	U	Chla	0.42 ug/l	SM 10200-H USGS I-4650-	24 hours
Leeds Point	Total Nitrogen	FwSw	U	TN	0.088 mg/l	0303 1-4650-	28 days
					<u> </u>	USGS I-4650-	,
Leeds Point	Total Phosphorus	FwSw	U	TP	0.025 mg/l	03	28 days
Leeds Point	Dissolved Ammonia	FwSw	F	DNH3	0.027 mg/l	350.1 MOD	14 days
Leeds Point	Dissolved Nitrite + Nitrate	FwSw	F	DNO3	0.027 mg/l	EPA 353.4	28 days
Leeds Point	Dissolved Orthophosphate	FwSw	F	DPO4	0.013 mg/l	EPA 365.5	28 days
Leeds Point	Dissolved Nitrogen	FwSw	F	DN	0.08815 mg/l	USGS I-4650- 03	28 days
LCCG3 i Ollit	Dissolved Nitrogen	1 WOW		DIV	0.02501	USGS I-4650-	20 day3
Leeds Point	Dissolved Phosphorus	FwSw	F	DP	mg/l	03	28 days
Leeds Point	Total Organic Nitrogen	FwSw	NA	TON	NA	Calculated	NA
Leeds Point	Dissolved Organic Nitrogen	FwSw	NA	DON	NA	Calculated	NA
Leeds Point	Dissolved Organic Phosphorus	FwSw	NA	DOP	NA	Calculated	NA
Leeds Point	Particulate Organic Nitrogen	FwSw	NA	PON	NA	Calculated	NA
Leeds Point	Particulate Phosphorus	FwSw	NA	PP	NA	Calculated	NA
OCUA	Biochemical Oxygen Demand (5 day)	FwSw	U	BOD5	1.0 mg/l	SM 5210B	48 hours
OCUA	Carbonaceous Oxygen Demand (5 day)	FwSw	U	CBOD5	1.0 mg/l	SM 5210B	48 hours
OCUA	Carbonaceous Oxygen Demand (20 day)	FwSw	U	CBOD20	1.0 mg/l	SM 5210B	48 hours
EPA Region 2	Alkalinity	FwSw	U	Alk	1.0 mg/l	SM 2320 B	14 days
EPA Region 2	Total Silica	FwSw	U	Si	1.0 mg/l	EPA 200.7 rev 4	6 months
EPA Region 2	Dissolved Silica	FwSw	F	DSi	1.0 mg/l	EPA 200.7 rev 4	6 months
EPA Region 2	Total Organic Carbon	Fw	U	TOC	1.0 mg/l	SM 5310 B	28 days
EPA Region 2	Dissolved Organic Carbon	Fw	F	DOC	1.0 mg/l	SM 5310 B	28 days
Leeds Point	Dissolved Inorganic Carbon	Sw	F	DIC	0.5 mg/l	SM 5310 C	28 days
Leeds Point	Dissolved Organic Carbon	Sw	F	DOC	0.5 mg/l	SM 5310 C	28 days
Leeds Point	Total Organic Carbon	Sw	F	TOC	0.5 mg/l	SM 5310 C	28 days
Leeds Point	Particulate Organic Carbon	Sw	NA	POC	NA	Calculated	NA

13. Chain Of Custody

Chain of custody procedures will be instituted for this project. Chain of custody procedures will be employed until samples reach the appropriate laboratory. Once samples reach the laboratory the laboratories internal sample tracking procedures will be utilized. (See Appendix B)

14. Corrective Action:

All laboratories involved in this project are required to maintain standard operating procedures which outline specific action to pursue should corrective action be necessary. If acceptable results cannot be obtained due to: either field or laboratory errors (calibration standards, proficiency testing samples, blanks, spikes, or duplicates falling out of range) the affected samples will be re-analyzed and steps will be taken to ensure that the data produced is accurate. Standards and reagents will be replaced, equipment will be checked, or other action, will be taken to remedy the situation. NJDEP designated project officers will be notified in writing anytime a deviation from the approved work plan has occurred.



Appendix A Sample Chain of Custody Form

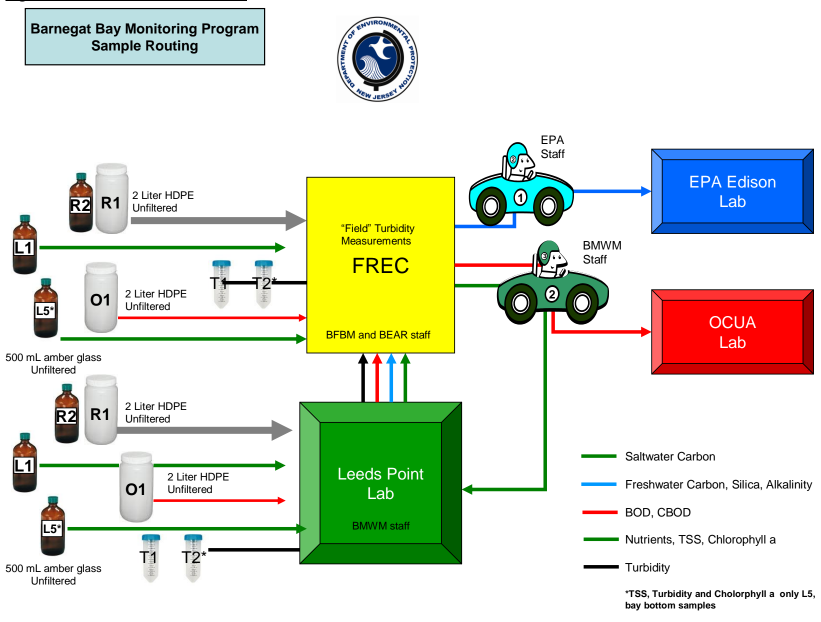
Appo	muix 1		Chain of Custody							
			General	Information		0				
Site #			Site Description			Sample # (Site # + Date)				
Sample Collection Date (mm/dd/yyyy)			Sample Collector							
						Sample Type (circle one)	В	imple lank		
Sample Collection Time (hh:mm)			Collector Organization			(* * * * * * * * * * * * * * * * * * *	Re	olicate		
				Surface	Bottom (Bay samples only)		Surface	Bottom (Bay samples only)		
Filtration Time (hh:mm)			Turbidity Measurement Time (hh:mm)		,,	Turbidity (NTU)				
(1111.11111)			Field Measuren	nents/Observations						
	Surface	Bottom (Bay only)		Surface	Bottom (Bay only)					
Field Measurement Time (hh:mm)			рН				Comments			
Sample Depth (ft)			Specific Conductance (circle units) uS/cm mS/cm							
Uncorrected Water Temp (deg C)			Ambient Transmissionmetry		NA					
Corrected Water Temperature (deg C)			Underwater Transmissionmetry		NA					
Dissolved Oxygen (mg/l)			Salinity (ppth)							
Dissolved Oxygen Saturation (%)	<u></u>		Secchi Depth (ft)		NA					
			Raw	Sample						
Container ID	C	ontainer	Matrix	Par	ameter	Fraction	Pres	ervative		
L1 - U	Amber I	HDPE, 500 mL	Freshwater / Saltwater	TSS, Chlorop	hyll a (bay only)	Total	Ice,	4 deg C		
O1 - U	Н	HDPE, 2L Freshwater / Saltwater		BOD5, CBOD5, CBOD20				Total	Ice, 4 deg C	
R1 - U		DPE, 2L DPE, 500 mL or	Freshwater / Saltwater			Total	Ice,	4 deg C		
R2 - UB	Н	IPDE,2L	Saltwater	TN, TP, Turbidity	, TOC, Silica Bottom	Total	Ice, 4 deg C			
L5 - UB		HDPE, 500 mL	Saltwater		a (bay only) Bottom	Total	Ice, 4 deg C			
L6 - U	Polyeth	nylene , 2 X 1L	Saltwater		plankton	Total	otal 0.5 % (v/v) glutaraldehy			
Container ID	C	ontainer	NJDEP Leeds Point Laborato Matrix		on #: 01179) ameter	Fraction	Pres	ervative		
L1 - U		HDPE, 500 mL	Freshwater / Saltwater		hyll a (bay only)	Total		4 deg C		
L2 - U		PE centrifuge tube	Freshwater / Saltwater		N, TP	Total		4 deg C		
L3 - F		PE centrifuge tube	Freshwater / Saltwater	DNH3		Dissolved		5% Phenol		
L4 - F	50 mL HDF	PE centrifuge tube	Freshwater / Saltwater	DNO3, DI	DNO3, DPO4, DN, DP		Ice,	4 deg C		
L5 - U	Amber I	HDPE, 500 mL	Saltwater	TSS, Chlorophyll	a (bay only) Bottom	Total	Ice,	4 deg C		
L6 - U	Polyeth	ylene , 2 X 1 L	Saltwater	Phyto	plankton	Total	0.5 % (v/v) gli	utaraldehyde, Ice		
L7 - U		ss, 250 mL	Saltwater		TOC	Total		2SO ₄ pH<2		
L8 - F L9-UB		ss, 250 mL	Saltwater		000	Dissolved Total		₂ SO ₄ pH<2 4 deg C		
L9-UB		PE centrifuge tube ss, 250 mL	Saltwater Saltwater		P Bottom Bottom	Total		4 deg C ₂SO ₄ pH<2		
			Ocean County Utilities Authority La							
Container ID	C	ontainer	Matrix	Par	ameter	Fraction	Pres	ervative		
O1 - U	H	DPE 2 L	Freshwater / Saltwater	BOD5, CB0	OD5, CBOD20	Total	Ice,	4 deg C		
			USEPA Region 2 Laboratory	at Edison Environme	ental Center					
Container ID	C	ontainer	Matrix		ameter	Fraction	Pres	ervative		
Container ID E1 - U		ontainer PE, 250 mL		Par		Fraction Total		ervative 4 deg C		
	HDF		Matrix	Par. Total	ameter		Ice,			
E1 - U	HDF	PE, 250 mL	Matrix Freshwater / Saltwater	Par Total S	ameter Alkalinity	Total	Ice,	4 deg C		
E1 - U E2 - U	HDF HDF	PE, 250 mL PE, 125 mL	Matrix Freshwater / Saltwater Freshwater / Saltwater	Par Total S	ameter Alkalinity ilica	Total Total	Ice, conc H	4 deg C NO₃ pH<2		
E1 - U E2 - U E3 - F	HDF HDF Glas	PE, 250 mL PE, 125 mL PE, 125 mL	Matrix Freshwater / Saltwater Freshwater / Saltwater Freshwater / Saltwater	Par Total S S	ameter Alkalinity illica	Total Total Dissolved	Ice, conc H conc H	4 deg C NO ₃ pH<2 NO ₃ pH<2		
E1 - U E2 - U E3 - F E4 - U	HDF HDF HDF Gla:	PE, 250 mL PE, 125 mL PE, 125 mL ss, 250 mL	Matrix Freshwater / Saltwater Freshwater / Saltwater Freshwater / Saltwater Freshwater Freshwater Saltwater	Par Total \$ \$ Total	ameter Alkalinity iilica iilica	Total Total Dissolved Total	Ice, conc H conc H conc H	4 deg C NO ₃ pH<2 NO ₃ pH<2 ₂ SO ₄ pH<2		
E1 - U E2 - U E3 - F E4 - U E5 - F E6-UB	HDF	PE, 250 mL PE, 125 mL PE, 125 mL ss, 250 mL ss, 250 mL PE, 125 mL	Matrix Freshwater / Saltwater Freshwater / Saltwater Freshwater / Saltwater Freshwater Freshwater Saltwater Chain	Par Total S S T E Silice	ameter Alkalinity ilica ilica OCC DOC Bottom	Total Total Dissolved Total Dissolved	Ice, conc H conc H conc H	4 deg C NO ₃ pH<2 NO ₃ pH<2 2SO ₄ pH<2 2SO ₄ pH<2		
E1 - U E2 - U E3 - F E4 - U E5 - F E6-UB Container ID .1-U, R1-U, O1-U, (L5-	HDF	PE, 250 mL PE, 125 mL PE, 125 mL ss, 250 mL ss, 250 mL	Matrix Freshwater / Saltwater Freshwater / Saltwater Freshwater / Saltwater Freshwater Freshwater Saltwater	Par Total S S T E Silice	Alkalinity iilica iilica OC OC DOC Bottom	Total Total Dissolved Total Dissolved	conc H conc H conc H conc H	4 deg C NO ₃ pH<2 NO ₃ pH<2 NO ₃ pH<2 SO ₄ pH<2 SO ₄ pH<2 SO ₄ pH<2		
E1 - U E2 - U E3 - F E4 - U E5 - F E6-UB Container ID 1-U, R1-U, O1-U, (L5-U, L6-U))	HDF	PE, 250 mL PE, 125 mL PE, 125 mL ss, 250 mL ss, 250 mL PE, 125 mL	Matrix Freshwater / Saltwater Freshwater / Saltwater Freshwater / Saltwater Freshwater Freshwater Saltwater Chain	Par Total S S S S Of Cusody Dat / /201	Alkalinity ilica ilica OCC OCC Bottom	Total Total Dissolved Total Dissolved Total Sample drop off a	conc H conc H conc H conc H conc H conc H	4 deg C NO ₃ pH<2 NO ₃ pH<2 SO ₄ pH<2 SO ₄ pH<2 SO ₄ pH<2 SO ₄ pH<2 FREC G		
E1 - U E2 - U E3 - F E4 - U E5 - F E6-UB Container ID .1-U, R1-U, O1-U, (L5-	HDF	PE, 250 mL PE, 125 mL PE, 125 mL ss, 250 mL ss, 250 mL PE, 125 mL	Matrix Freshwater / Saltwater Freshwater / Saltwater Freshwater / Saltwater Freshwater Freshwater Saltwater Chain	Par Total S S S T C Silica of Cusody	ameter Alkalinity iilica iilica OC OC IBottom	Total Total Dissolved Total Dissolved Total Sample drop off a	conc H conc H conc H conc H	4 deg C NO ₃ pH<2 NO ₃ pH<2 SO ₄ pH<2 SO ₄ pH<2 SO ₄ pH<2 SO ₄ pH<2 FREC c		

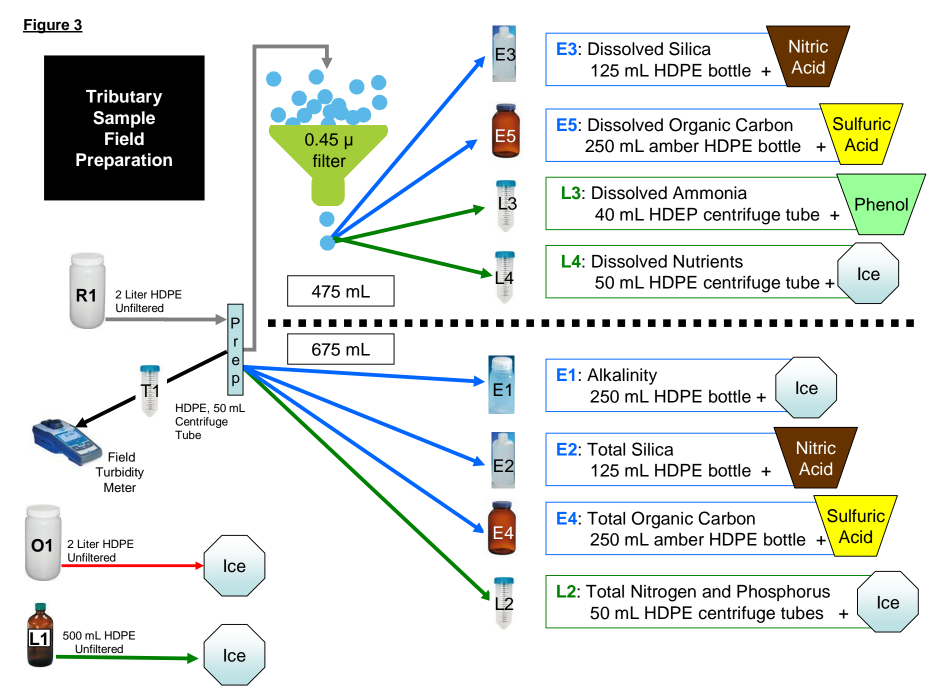
Revision 2, December 2011 20

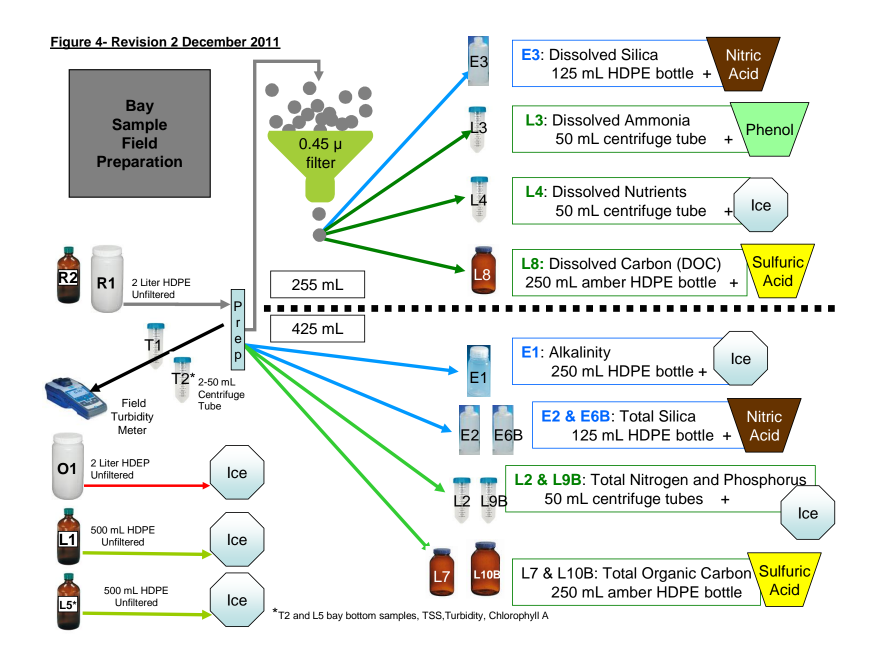
Appendix B Sample Routing and Field Preparation Figures 2-4

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Figure 2- Revision 2 December 2011







Appendix C: Example Field Calibration

Field Calibration Record-Barnegat Bay Monitoring Project

<u>pH Meter Calibration</u> Required Accuracy: \pm 0.05 standard units (su) of the true value for the calibration buffers used, \pm 0.10 for the mid range calibration check buffer. The temperature of the buffers must be recorded (if available).
Buffer Temp "As Found" Meter Reading Set Meter Reading Time Date Tech
4.00
7.00
10.00
Buffer used for Calibration Check:Temp of Buffer:
Calibration Check Buffer result (performed in measure mode):
Time of Calibration Check and Tech:
Conductivity Meter Calibration Required Accuracy: Within 1% of the true value for the standard used. Meter is calibrated according to manufacturer's instructions. Standard check is
required each day of use.
mS/cm Standard used for Calibration Check (performed in measure mode):
Date of Calibration Check:
Tech:
DO Meter Calibration
Meters are to be calibrated each day of use against air or water saturated air.
Meters also require a Winkler test each week of testing.
Required Accuracy between the Winkler titration and meter: \pm 0.3 mg/L
Normality of Titrant (from container of sodium thiosulfate):
Beginning mls:
Ending mls:
Change in mls:
DO reading from Meter for the sample used for Winkler:
Tech:
Date:
Temperature Calibration
Thermometers are due for the next calibration on August 17, 2011 or prior to use
for any sampling events just prior to that date. Calibration performed on May 17

2011 expires on August 17, 2011. Records of the calibration will be provided by the

person calibrating the thermometer if done by DEP staff.

daw 6/22/11

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