0600Bm--Jack Fitz Cruise 02 May 21-31 2010

*****DATA SOURCE*****
Data were compiled from surveys conducted in the Gulf of Mexico.

Data were compiled from NewFields Environmental Forensics Practice, LLC
(Alpha) lab electronic data from QC Batches 1005070, 1005071, 1005072,
1005077, 1005079, 1005083, 1005084, 1006018, 1006020, 1006021, 1006022,
1006024, 1006025, 1006026, 1006027, 1006028, 1006029, and 1006030.

*****DATA COLLECTION PURPOSE*****
Natural Resource Damage Assessment

*****DATA USE QUALIFICATION*****
These data are a subset of samples collected on Jack Fitz Cruise 2. As
more data become available, they will be added to this data set.

*****STUDY*****
The data include water and oil chemistry data. Oil chemistry data are
stored in the samptar/chemtar tables, and are reported with solid units.

*****STATION*****
StationIDs are based on the Grid locations recorded in the NOAA Field
Sampling Information database, plus a sequential number used for each
distinct latitude/longitude position reported. Datum was not provided
but assumed to be NAD83.

*****SAMPLES AND REPLICATES*****
The original SampleIDs reported by the lab from the Chain-of-Custody is
stored in the ExSampID field. Whole water samples noted with a *ww* as
part of the FldSampID, and VOA samples (unfiltered, with a *wv* as part
of the FldSampID) are represented with matrix of "WH." Filtered
(dissolved) samples were noted with a 'wd' in the original sampleID, a
"F" at the end of the Query Manager SampleID, and coded with a Matrix of
"DS." Particulate samples are noted with a *wp* in the original sampleID,
coded with a "P" at the end of the sampleID, and a Matrix of "PT." Note
that the particulate fraction represents the filter that has been
analyzed after flushing with a volume of water; thus the concentrations
are provided in a liquid basis.

The collection depth of water samples in the fields UDepth and LDepth are
reported in meters.

The default labrep code "1A" was used for most data. Lab duplicates
are noted with a "2" as the first letter of the labrep.

Several analytes are reported from 2 different analytical methods. The
"preferred" result (usually with lower detection limits) is given the
default labrep code (e.g., "1A" or "2A"). The results from the non-
preferred analytical method have a "X" appended to the labrep code (e.g.,
"1AX" or "2AX")

The following chemcode/analytes were measured using two methods:
Methods: PIANO Volatile Hydrocarbons by GC/MS | 8260M and Total Saturated Hydrocarbons by GC/FID | 8015M
AHCN_C09/ Nonane
AHCN_C10/ Decane
AHCN_C11/ Undecane
AHCN_C12/ Dodecane
AHCN_C13/ Tridecane
The results for PIANO Volatile Hydrocarbons by GC/MS were assigned labrep "1AX"

Methods: PIANO Volatile Hydrocarbons by GC/MS | 8260M and Alkylated Polynuclear Aromatic Hydrocarbons | 8270M
BTHIOPHNE/ Benzo(b)thiophene
METHNAP_1/ 1-Methylnaphthalene
METHNAP_2/ 2-Methylnaphthalene
NAPTHALENE/ Naphthalene
The results for PIANO Volatile Hydrocarbons by GC/MS were assigned labrep "1AX"

Alpha Lab Analytical Methods:
Total Saturated Hydrocarbons by GC/FID | 8015M | SOP. 0-003 Rev. 5
(abbreviated as 8015 M - Tot Sat. HC - GC/FID)
Alkylated Polynuclear Aromatic Hydrocarbons | 8270M | SOP. 0-008 Rev. 6
(abbreviated as 8270 M - Alkylated PAHs)
PIANO Volatile Hydrocarbons by GC/MS | 8260M | SOP. 0-019 Rev. 2
(abbreviated as 8260 M - PIANO VolHC - GC/MS)

****SUMMED PARAMETERS****
No sums were calculated.

****QUALIFIERS****
Qualifiers recorded in the chemistry files represent the final data qualifiers provided by the data validation. Descriptions of the data qualifiers are included in the data dictionary.

****OTHER****
The original analyte reported as Benzo(k)fluoranthene was identified by the data validators to be a coelution of Benzo(k)fluoranthene and Benzo(j)fluoranthene. Therefore, the chemical data for the original Benzo(k)fluoranthene results have been assigned a chemical code for Benzo(j+k)fluoranthene.