

Abbreviations

	SeaBASS and current report abbreviation	SeaHARRE and/or previous report abbreviation	full name	notes
Primary Pigments	Allo	Allo	alloxanthin	
	alpha-beta-Car	Caro	carotenes	alpha (beta, epsilon) + beta (beta, beta) carotene. Unresolved and therefore undifferentiated
	But-fuco Diadino Diato	But fuco Diad Diato	19'-butanoyloxyfucoxanthin diadinoxanthin diatoxanthin	
	Fuco Hex-fuco	Fuco Hex fuco	fucoxanthin 19'-hexanoyloxyfucoxanthin	
	Perid	Perid	Peridinin	
	Tot_ChI_a	TChI a	total chlorophyll a	DV_ChI_a + MV_ChI_a + Chlide_a + ChI_a allomers + ChI_a epimers
	Tot_ChI_b Tot_ChI_c Zea	TChI b TChI c Zea	total chlorophyll b total chlorophyll c Zeaxanthin	DV_ChI_b + MV_ChI_b + ChI_b epimers ChI_c3 + ChI_c12

Secondary

Pigments

Chl_c3	Chl c3	Chlorophyll c3
Chlide_a	Chlide a	chlorophyllide a
DV_ChI_a	DVChI a	divinyl chlorophyll a
DV_ChI_b	DVChI b	divinyl chlorophyll b
MV_ChI_a	ChI a	monovinyl chlorophyll a
MV_ChI_b	ChI b	monovinyl chlorophyll b
		Chlorophyll c2 + chlorophyll c1 +
	ChI c12	MGDVP
		Mg-2,4-divinyl pheoporphyrin a5
	MGDVP	monomethyl ester

Tertiary

Lut	Lut	Lutein	
Neo	Neo	Neoxanthin	
Phide_a	Phide a	total pheophorbide a	multiple peaks
Phytin_a	Phytin a	total pheophytin a	pheophytin a + pheophytin a'
Pras	Pras	Prasinoxanthin	
Viola	Viola	Violaxanthin	

Ancillary

Pigment	Gyro	Gyr diester	Gyroxanthin diester
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Replicate filters

The replicate filter precision page summarizes our results for any replicate filters you submitted. On both the replicate filter and analysis precision page, pairs with precision worse than 10% (15% for degradation products) are flagged in yellow. If a simple reason can be determined (ex. Concentration is below the effective LOQ), it is noted in a comment.

Replicate injections

The analysis precision page summarizes our results for the same sample extract injected twice. Typically, we reinject the first sample analyzed on a given at the end of the day (the ".5" injection). For example, sample 03-0001 and 03-0001.5 are replicate injections of the same extract, injected approximately 24 hours apart (all samples extracted on a particular day require about 24 hours to complete the HPLC analyses). We do this to measure our analysis precision and any effects caused by a sample's residence time in the refrigerated autosampler compartment. Please note that individual results with very large CV% are usually caused by pigments present in very low concentrations. On both the replicate filter and analysis precision page, pairs with precision worse than 10% (15% for degradation products) are flagged in yellow. If a simple reason can be determined (ex. Concentration is below the effective LOQ), it is noted in a comment.

Effective Limit of Quantitation

On the effective LOQ page, we calculate an effective limit of quantitation based on our calculated LOQs (calculated in ng/injection), our typical extraction volume for this sample set, and the various filtration volumes used with your samples. We make these calculations because our LOQ information is most useful to the data user if it is available in units of concentration (ug/L seawater). The same LOQ can end up looking very different for different filtration volumes. For example, the LOQ of 0.25 ng will result in very different effective LOQs when carried through our calculation equation to represent the ug/L seawater. For an extraction volume of 2.5 ml and a filtration volume of 2800 ml, the calculated effective LOQ would be 0.002 ug/L. However, if the filtration volume were only 100 ml, the effective LOQ would calculate to be 0.042 ug/L. Without these calculations, the end user has no way of knowing that both of these concentrations were acquired at detection-limited concentrations.

Zeros

Instead of including zeros, pigments that were "not found" (not detected) are noted with a replacement value of **-8888**. For pigments that have a replacement value in the respective cell, the pigment was investigated and determined to be

Analysis method description

The HPLC analysis method can be cited as Van Heukelem and Thomas (2001), further described in Hooker et al. (2005). For a more detailed description, please see below; contact Crystal for a tailored description.

The HPLC used for pigment analysis is an Agilent RR1200 with a programmable autoinjector (900 ul syringe head), refrigerated autosampler compartment, thermostatted column compartment, quaternary pump with in-line vacuum degasser, and photo-diode array detector with deuterium and tungsten lamps. The HPLC is controlled by Agilent Chemstation software.

The 4.6 x 150 mm HPLC Eclipse XDB column (Agilent Technologies, Palo Alto, CA) is filled with a C8 stationary phase (3.5 um stationary phase); the mobile phase consists of a linear gradient from 5-95% solvent B over

27 minutes, for which solvent A is 70 parts methanol, 30 parts 28 mM tetrabutylammonium acetate (pH 6.5) and solvent B is methanol. The column temperature is 60 C and the photo diode array detector is set to plot chromatograms at 450, 665, and 222 nm to acquire visible absorbance spectra between 350 and 750 nm.

Vitamin E acetate is used as the internal standard (ISTD) for determining extraction volumes. Its absorbance is monitored at 222 nm; it has negligible absorbance at 450 nm and none at 665 nm. Therefore, it does not interfere at wavelengths used to quantify pigments and can be used in very high concentrations with S:N ratios much higher than are possible with pigments. The high signal:noise ratio contributes to excellent analysis precision, for which

injection repeatability averages 0.6%. It is stable under conditions of extraction and analysis.

Calibration is performed with individual pigment standards, whose concentrations have been determined spectrophotometrically using absorption coefficients in common with those used by most other laboratories (Hooker et al. 2005) and the commercial vendor, DHI Water and Environment (Horsholm, Denmark). Standards are either purchased from DHI (in solution with concentrations provided) or purchased in solid form and suspended in solvent at GSFC.

Thirty-six peaks are individually quantified by HPLC, from which 26 pigments are reported (some pigments contain individual components that are summed and reported as one pigment).

SeaBASS submission

Please refer to the "Data Contributors" menu on the SeaBASS website (<http://seabass.gsfc.nasa.gov>) for information on how to prepare your data files for submission. If your data file contains measurements that were below detection limits ("not found", see Zeros section), those values were set to -111, and the following information should be included in your metadata headers:

/below_detection_limit=-111

! Comments

!

! Measurements below detection limits are assigned the value -111

SeaBASS

abbreviation	description	notes
DP	total diagnostic pigments	PSC + allo + zea + Tot_ChI_b
PPC	photoprotective carotenoids	allo + diadino + diato + zea + alpha-beta-car
PPC_TCar	ratio of photoprotective carotenoids to total carotenoids	[PPC]/[Tcar]
PPC_TPg	ratio of photoprotective carotenoids to total pigments	[PPC]/[TPg]
PSC	photosynthetic carotenoids	but-fuco + fuco + hex-fuco + perid
PSC_TCar	ratio of photosynthetic carotenoids to total carotenoids	[PSC]/[TCar]
PSP	photosynthetic pigments	PSC + TChI
PSP_TPg	ratio of photosynthetic pigments to total pigments	[PSP]/[TPg]
TAcc	total accessory pigments	PPC + PSC + Tot_ChI_b + Tot_ChI_c
TAcc_TChIa	ratio of total accessory pigments to total chlorophyll a	[Tacc]/[TchIa]
TCar	total carotenoids	PPC + PSC
TChI	total chlorophylls	Tot_ChI_a + Tot_ChI_b + Tot_ChI_c
TChI_TCar	ratio of total chlorophyll to total carotenoids	[TChI]/[TCaro]
TChIa_TPg	ratio of total chlorophyll a to total pigments	[TChIa]/[TPg]

TPg

total pigments

TAcc + Tot_ChI_a