

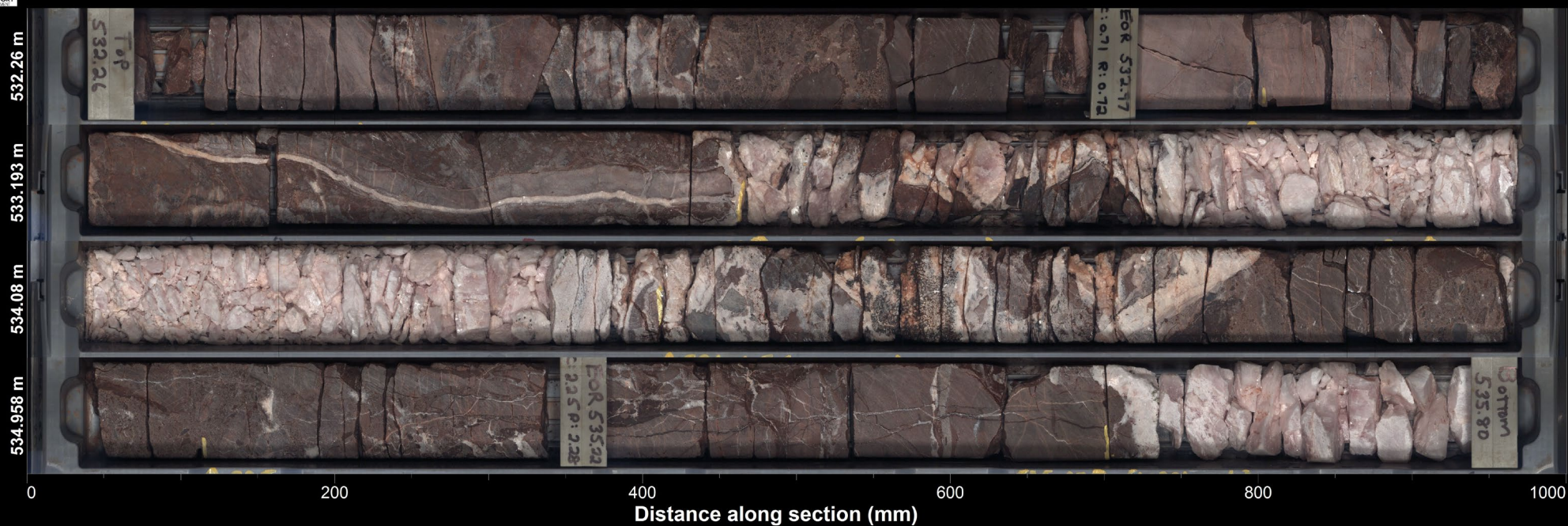
## HyLogger Data Package 0074

HyLogger drillhole report for Manbulloo S1,  
greater McArthur Basin, Northern Territory.

Belinda Smith



NVCL NTGS Node - 8446303\_Manbulloo\_S1 Tray 0034, 532.3 to 535.8 m



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## The Spectral Geologist Advisory

The results in this report were obtained using The Spectral Geologist (TSG) software. This software uses The Spectral Assistant (TSA) as a default to identify minerals and their abundances for the Short Wave Infrared (SWIR) and Thermal Infrared (TIR) spectrum. TSA is a general unmixing algorithm and is trained on a relatively small subset of commonly-occurring minerals. It does not make the right identifications all of the time. The unmixing is an interpretation result of 'best fit'. TSA abundances are relative abundances, only the two (or three) most spectrally active minerals identified in the SWIR and the three (sometimes four) most spectrally active minerals in the TIR wavelengths are reported. If there are more than two or three minerals actually present in the sample in the SWIR (or three to four minerals in the TIR) then this is not reflected AT ALL in the reported abundances. Minerals are reported as a fraction of the overall spectral fit rather than actual quantifiable concentrations (total minerals present add up to 1). The SWIR wavelength only identifies hydrous silicates and carbonates. It does not reflect the TOTAL mineralogy of the sample. NTGS processed datasets exclude some minerals in the TSA library if the mineral is a poor spectral fit or unlikely in that geological environment, introducing a further element of interpretation.

Since April 2014, the TIR spectral responses have also been matched to minerals using Constrained Least Squares (CLS), which is an alternative unmixing classifier. CLS uses a Restricted Mineral Set (RMS) to minimise non-unique mineral modelling in the TIR spectrum. The RMS is determined by the processor who interprets 'domains' (hole intervals interpreted to have similar mineralogy) and then limits the set of possible mineral matches based on the geological understanding and spectral characteristics of that domain.

Since April 2017, the TIR spectral responses may also be matched to minerals using joint Constrained Least Squares (jCLST), which is an unmixing classifier that replaces the earlier system TSA (sTSAT) used in TSG versions 7 or earlier. In TSG8, jCLST is the default system unmixing algorithm, which interprets the TIR data using the results from the SWIR spectra and using scalars focussing on selected features in the Visible Near Infrared (VNIR) and TIR wavelengths. TSG8 datasets may have TIR mineral results reported as domained TSA (dTSAT), user TSA (uTSAT) or domained CLS (TIR-CLS1).

Any results from the TIR should be used with caution as algorithms and TSA libraries are in a constant state of revision. More information about the samples in the TIR reference library can be found in Schodlok *et al* (2016a).

These results were published using TSG Version 8.0.3.16 dated December 2018.

Please note: the results in this report are an interpretation from the spectral response.

These notes are designed to be read in conjunction with navigating through the TSG dataset (the TSG dataset is part of the HyLogger Data Package); the following pages are saved screens images within the \*holename\_tsg.tsg file. Navigate to the screens by using the View | Plot Layout pathway shown at the top right of each page. It is recommended to have this document open on one screen and the TSG dataset on the other.



## Manbulloo S1: Introduction

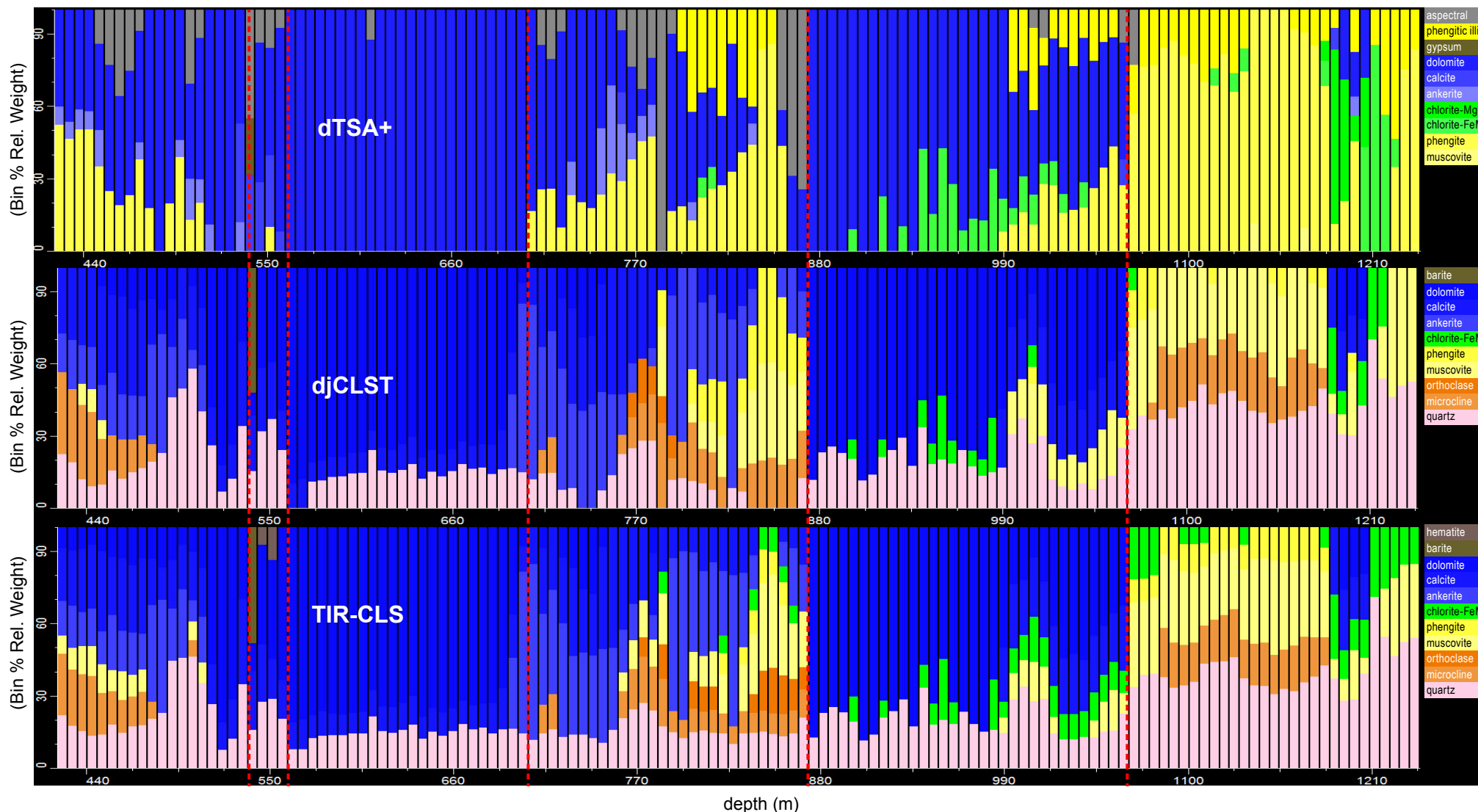
<b>Hole ID</b>	Manbulloo S1	<b>Unique identifier</b>	8446303
<b>Geological terrane</b>	McArthur Basin	<b>Total depth</b>	1236.22 m (from derrick floor)
<b>Latitude GDA94</b>	-14.927674°	<b>Longitude GDA94</b>	132.268419°
<b>Easting MGA94</b>	608 512 (Zone 53)	<b>Northing MGA94</b>	8 226 502 (Zone 53)
<b>Dip</b>	-90°	<b>Azimuth</b>	360°
<b>Logged by</b>	Pangaea Resources Pty Ltd	<b>Logged report ref</b>	Pangaea (2015): PR2015-0017
<b>Start core depth</b>	422.7 m (from derrick floor)	<b>End core depth</b>	1236.22 m
<b>Date HyLogged</b>	September 2014	<b>HyLogged by</b>	Darren Bowbridge
<b>Date of HyLogger report</b>	December 2018	<b>HyLogger report author</b>	Belinda Smith
<b>TSG version and build</b>	8.0.2.17 (December 2018)	<b>TSG product level</b>	3 (Huntington 2010)

Summary of information from Pangaea (2015):

- Drilled as a petroleum stratigraphic core well. Note the depths on the core trays (and from that, the HyLogger depths) are for depths from the Derrick Floor ('Driller depths'). This is 1.3 m higher than collar depth. Note this for any xyz coordinate generation from ground level.
- Coring commenced in the 'undifferentiated upper McArthur'. The logged stratigraphy notes that below the undifferentiated upper McArthur' is the 'Reward Dolomite equivalent'. This may be the Killaloc Formation (?) as noted by Hallett (2018).
- Below the 'Reward Dolomite equivalent' is the Fraynes Formation. Pangaea noted that this is equivalent to the Barney Creek Formation.
- The cored interval passed through the Fraynes Formation, into the Campbell Springs Dolostone, Blue Hole Formation, Kunja Siltstone and Mallabah Dolostone before terminating in the Amos Knob Formation.
- The wireline logs (in Enclosure 1 of Pangaea 2015) record gamma ray measurements to ~907 m, not to EOH (1236 m). The compensated density and borehole corrected gamma data measure the drillers depth (depth from Derrick Floor), which is 1.3 m above ground level. These are the values used as the HyLogger depths are depths from Derrick Floor (to match the driller core block and core tray markings). These wireline log measurements were imported into the TSG dataset. The point depth values were given interval depths by adding half the distance to the adjacent depth measurement.
- Most of the drillhole was HyLogged in September 2014. A processed dataset was released dated September 2016. However, trays 34–40 were not part of this original scan or processed dataset. These trays were scanned in October 2017, appended to the dataset and reprocessed. The current dataset differs from this earlier version as it has the missing trays, as well as petrophysics measurements and wireline data imported within the TSG dataset.



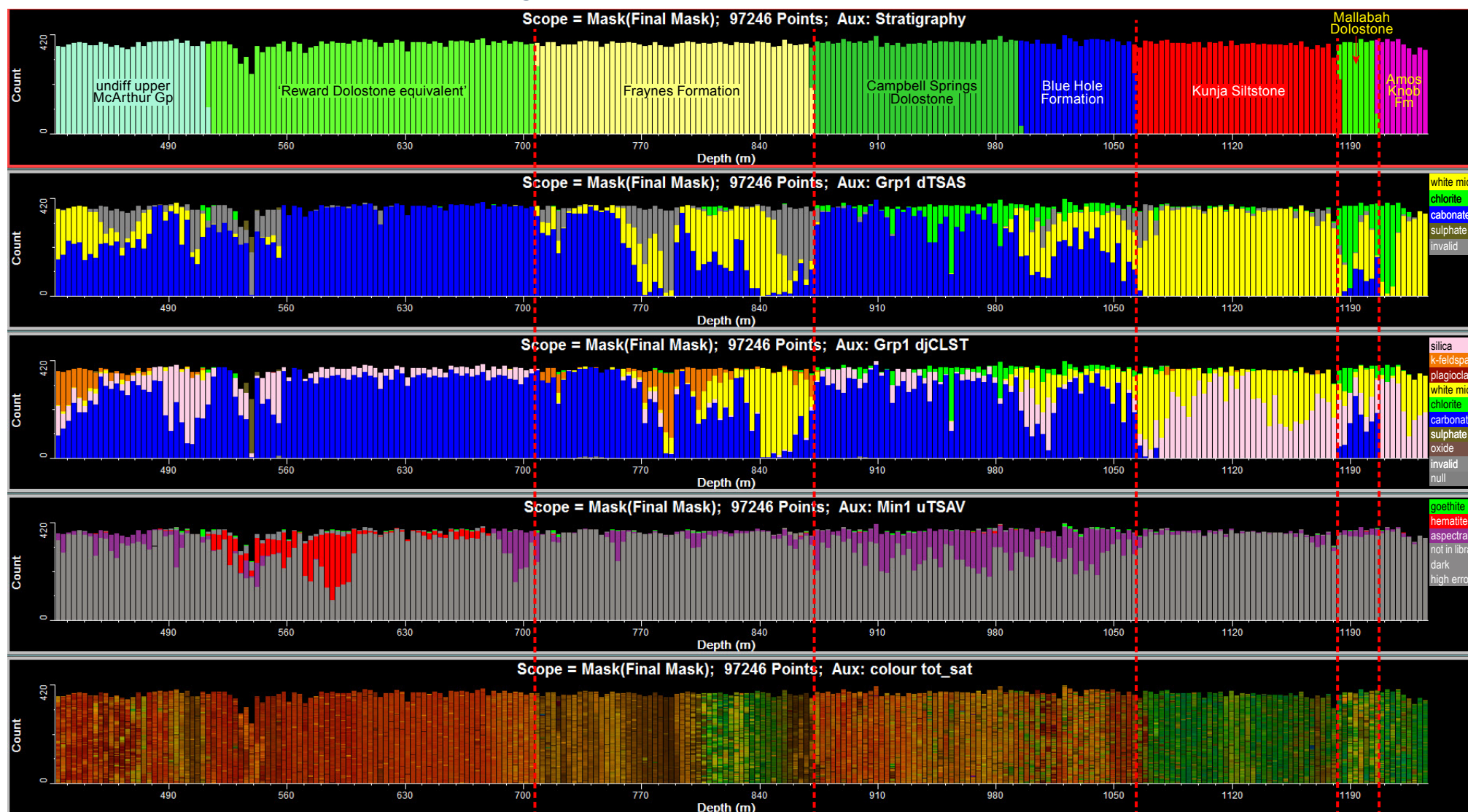
# Manbulloo S1: Mineral summary – all minerals, TSA and CLS



From Summary Screen: Row 1 is the domained SWIR results using TSA+. Row 2 is the TIR results derived from a domained joint CLS algorithm (see Guide to Scalars for a description of both TSA+ and djCLST). In this dataset, djCLST is used in preference over TIR-CLS1 and dTSAT (which was used in HDPs prior to HDP0067). Both TIR modelled results are shown here for comparison as the TIR spectra have non-unique results. Dotted lines delineate mineral changes, although there are smaller mineral changes not delineated here. There are sharply defined mineral changes around 870 m: from carbonate (blue) dominant to quartz dominant (with depth). Manbulloo S1 intersects a range of stratigraphic intervals, so the sharp mineral changes downhole should delineate those stratigraphic changes.

# Manbulloo S1: Mineral summary

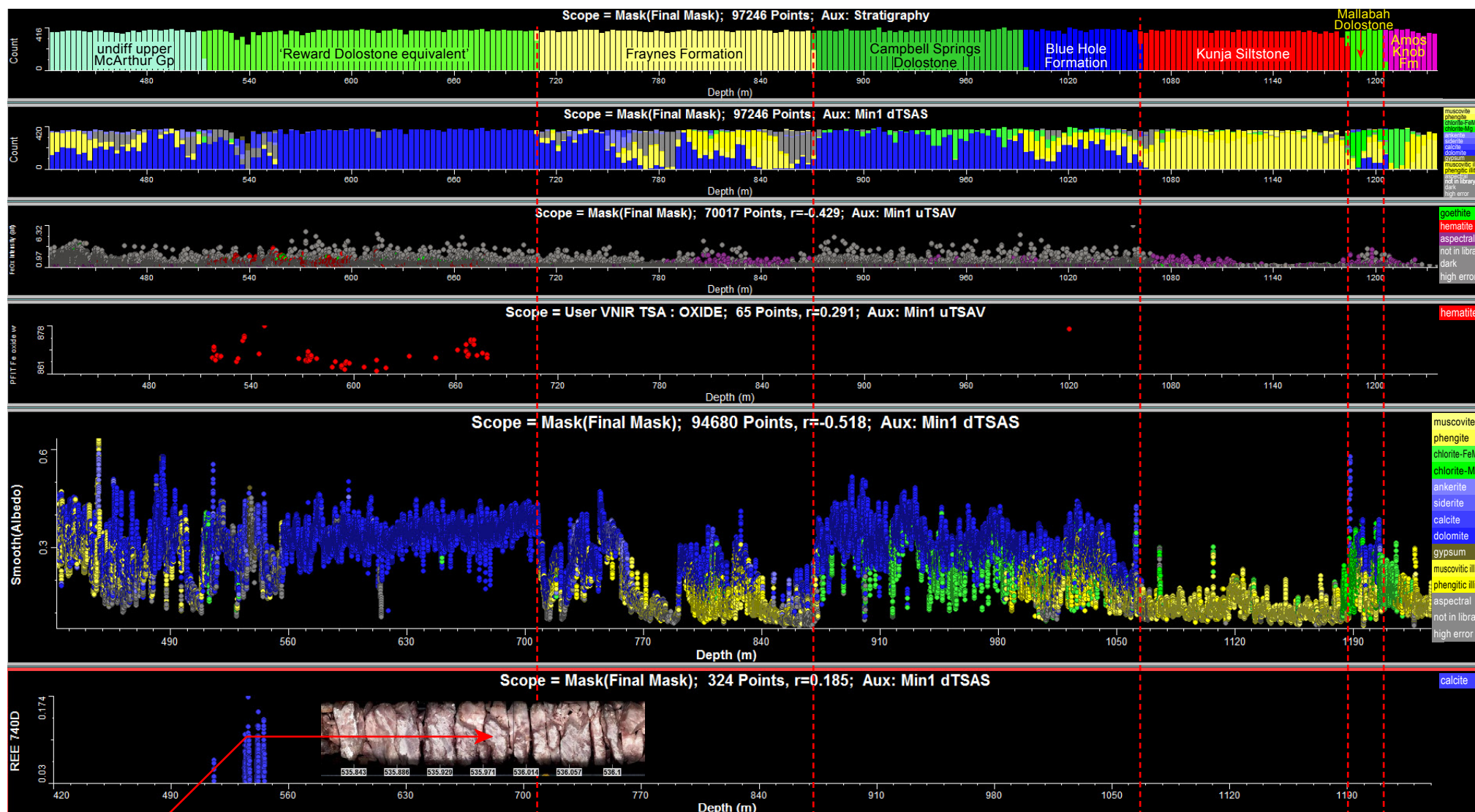
(View | Plot Layouts Load file 1\_VNIRSWIR: Mineral Summary)



Row 1 shows the logged stratigraphic intervals from Pangaea (2015). Rows 2 and 3 plot the SWIR and TIR spectra, coloured by the dominant mineral group match respectively. Row 4 plots the VNIR spectra, coloured to either goethite, hematite, spectral, or Not In Library (VNIR has a limited TSA mineral library for matching). Row 5 is the core colour. Dotted lines show mineral group (or core colour) boundaries. Most of the stratigraphic units in the lower part of Manbulloo S1 have sharply defined stratigraphic boundaries. The boundary between the Campbell Springs Dolostone and the Blue Hole Formation is the exception. The Fraynes Formation has several mineralogical and core colour changes. These may reflect depositional cycles (?). There is no clear mineralogical boundary between the upper 'Reward Dolostone equivalent' and the 'undifferentiated upper McArthur Group sediments'. However, within the 'Reward Dolostone equivalent', there is a change around 554 m. Below this depth; carbonate >> quartz. Above this depth, quartz is more common.

# Manbulloo S1: SWIR and VNIR mineral summary

(View | Plot Layouts Load file  
2\_VNIRSWIR: SWIR VNIR Summary)

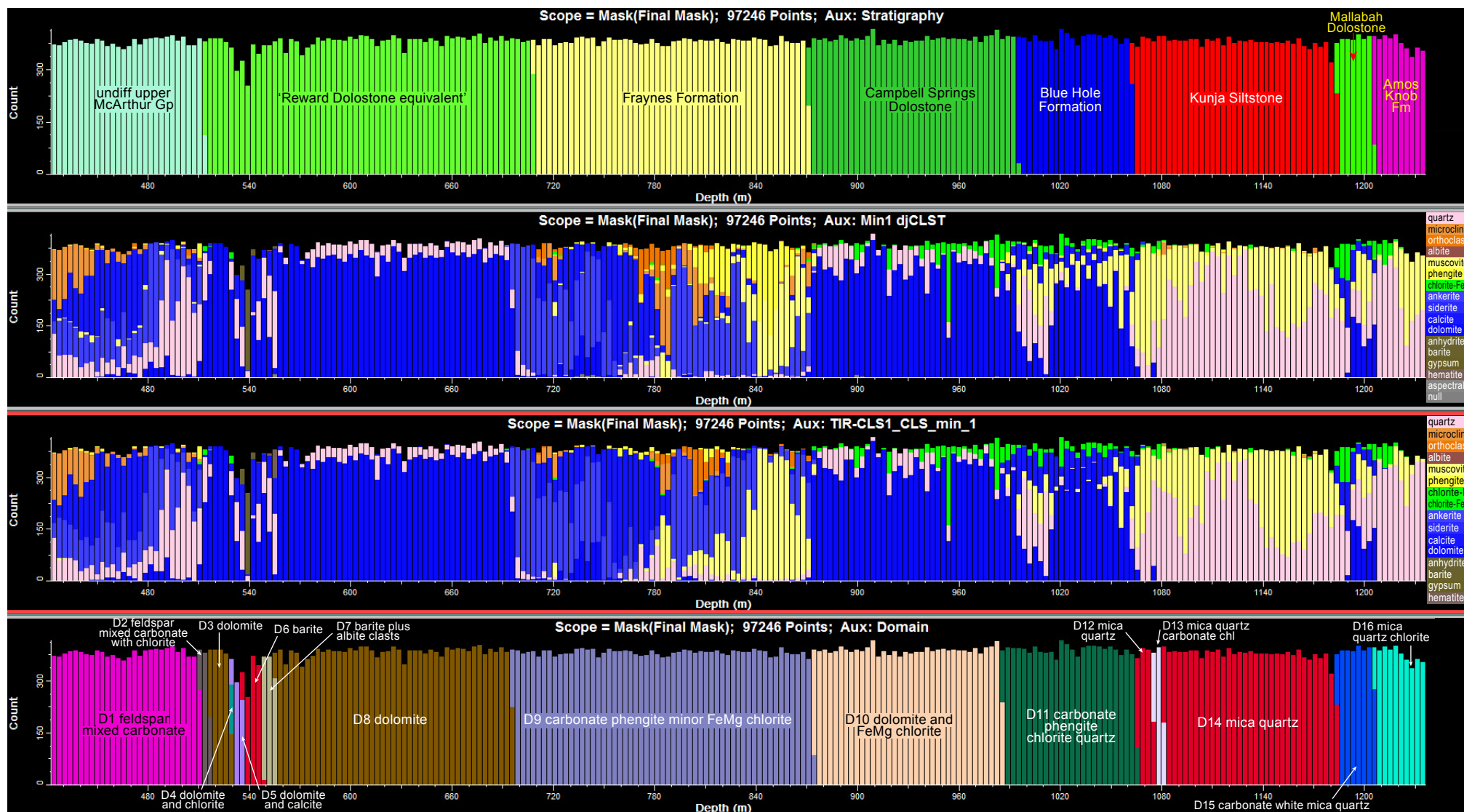


Row 1 shows the logged stratigraphic intervals from Pangaea (2015). Row 2 is the SWIR spectra coloured by the dominant mineral (may be in a mineral mixture). Row 3 is the VNIR spectra plotted against FeOx intensity batch scalar and coloured by the match to VNIR minerals. Row 4 are the VNIR spectra that match to either goethite or hematite, plotted by the wavelength of the Fe oxide feature. Hematite has a shorter wavelength feature compared with goethite. This scalar indicates hematite (but not goethite) is present 517–678 m. Row 5 plots the SWIR spectra by the smoothed albedo, coloured by the dominant SWIR mineral. Higher values are carbonates. Row 6 plots the spectra with a sharp 740 nm feature (see inset; left). This spectral response indicates the presence of Nd, a rare earth element (REE). The REE is discontinuously found within calcite between 515–546 m (see inset above). Dotted lines show albedo changes, which also indicate changes in carbonate content.



# Manbulloo S1: TIR mineral summary - overview

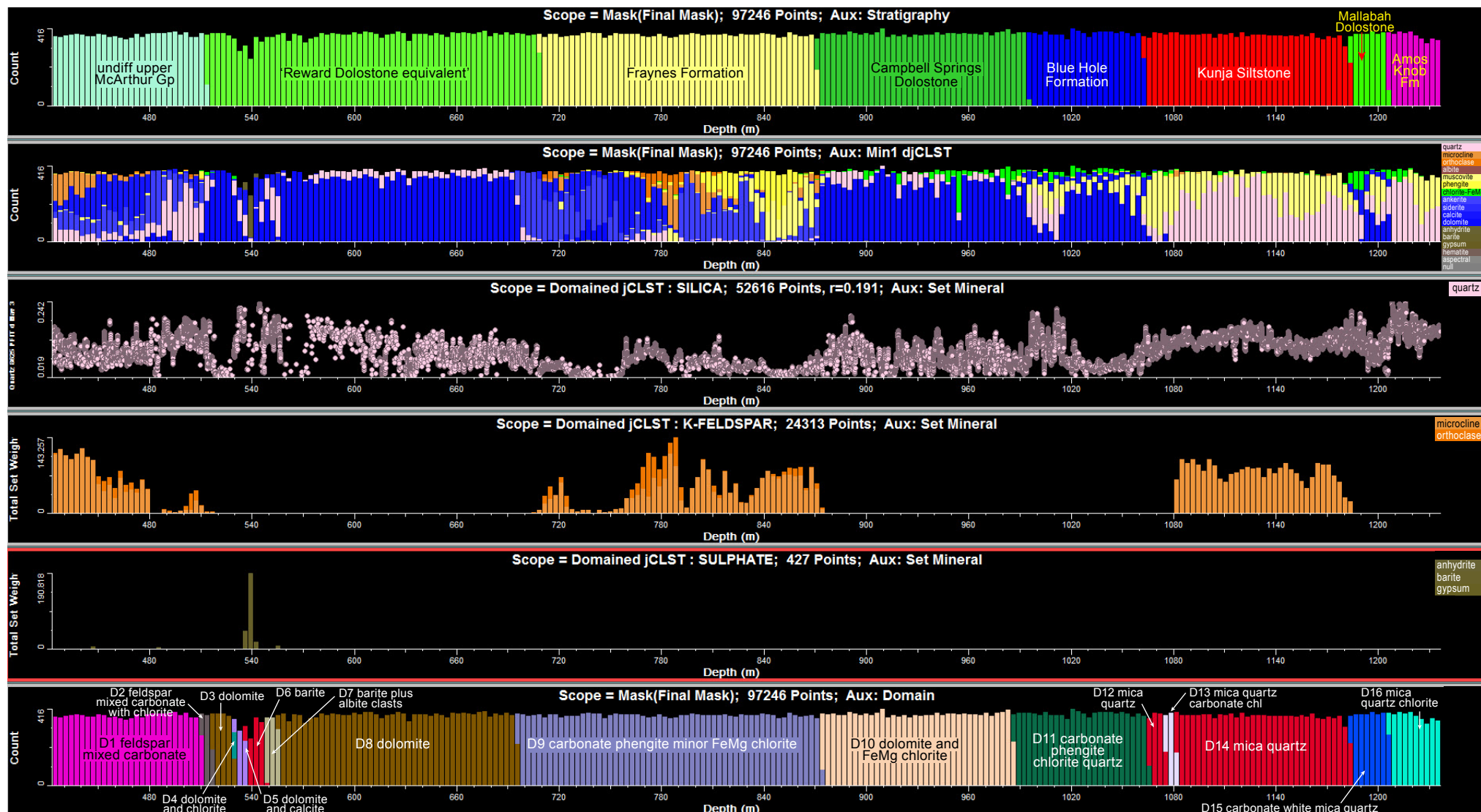
(View | Plot Layouts Load file 1\_TIR: TIR Summary)



Row 1 shows the logged stratigraphy from Pangaea (2015). Rows 2 and 3 plot the dominant TIR mineral (from a mineral mix) using a domained restricted mineral set (RMS). Row 2 uses the joint CLS algorithm that takes the results from a modified TSA and scalars operating on selected features in the VNIR and SWIR. Row 3 is the CLS algorithm that uses an RMS interpreted for each domain, allowing up to 6 mineral results per spectrum (dominant mineral shown here). In Manbulloo S1, the dominant TIR mineral result is similar from both algorithms. Row 4 is the domains interpreted for this hole. For more information on domains in the TSG dataset, open the 'D' tool on the toolbar. Note that the logged lithology within a stratigraphic interval can have different mineralogy. For example, within the upper part of the 'Reward Dolostone equivalent', there are several smaller intervals that delineate barite-bearing carbonate, plus smaller intervals of calcite and dolomite (little or no quartz).

# Manbulloo S1: TIR mineral summary 2

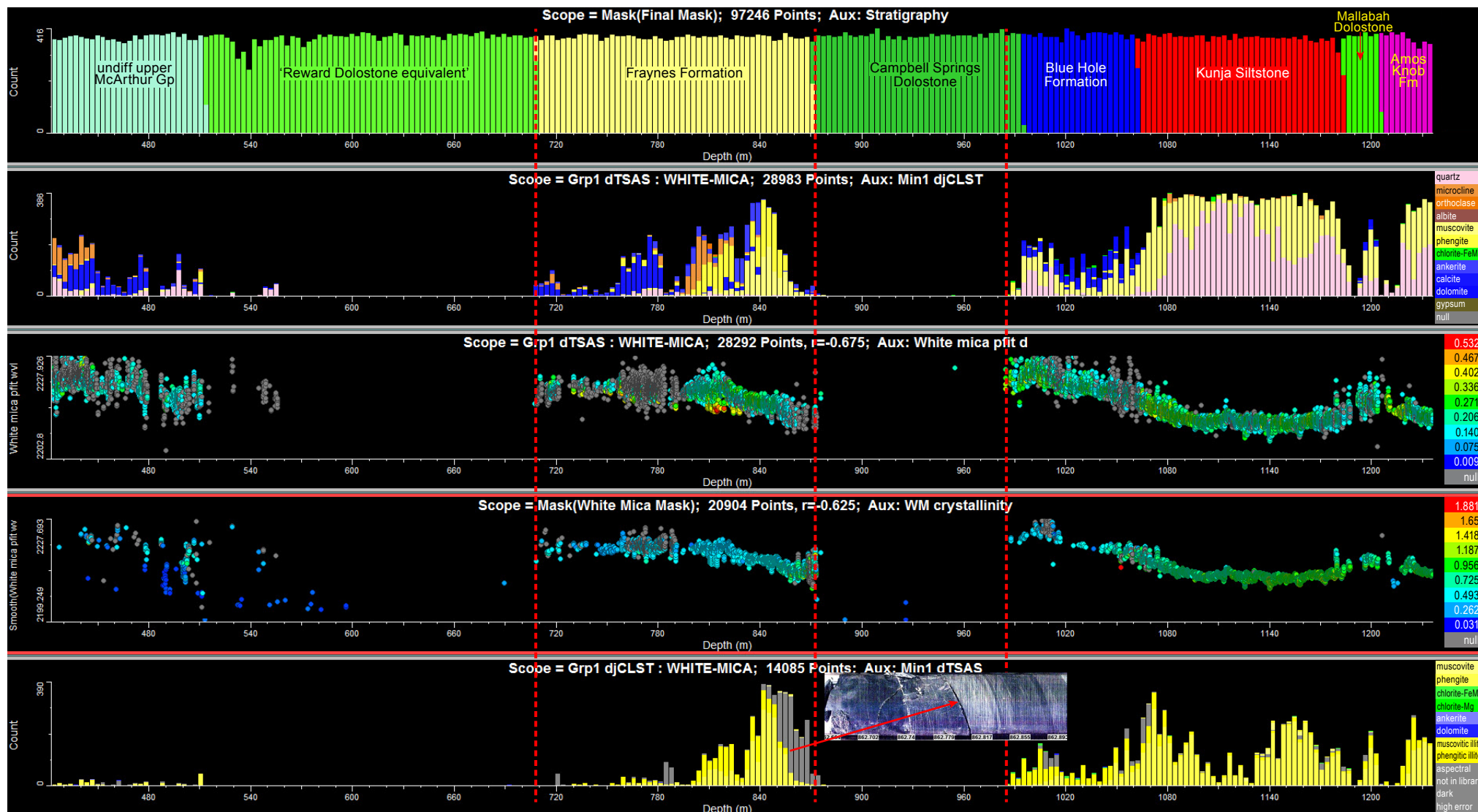
(View | Plot Layouts Load file1\_TIR: TIR Mineral Summary 2)



Row 1 shows the logged stratigraphic intervals from Pangaea (2015). Row 2 is the dominant TIR mineral from the domained joint CLS algorithm. Row 3 are the TIR spectra that contain a match to quartz, plotted by the depth of the quartz 8625 nm feature (analogous to abundance). Rows 4 and 5 plot the TIR spectra that contain a match to K-feldspar and sulphate respectively. K-feldspar is found grouped in 3 stratigraphic intervals: at the top of the cored interval (in the undifferentiated McArthur Group); within the Fraynes Formation; and throughout the Kunja Siltstone. The sulphate is dominantly coarse crystals of barite around 540 m. Row 6 is the interpreted domains, used for restricting the list of minerals that are used for mineral matching in the TIR.

# Manbulloo S1: White micas

(View | Plot Layouts Load file 3\_VNIRSWIR: White Micas)

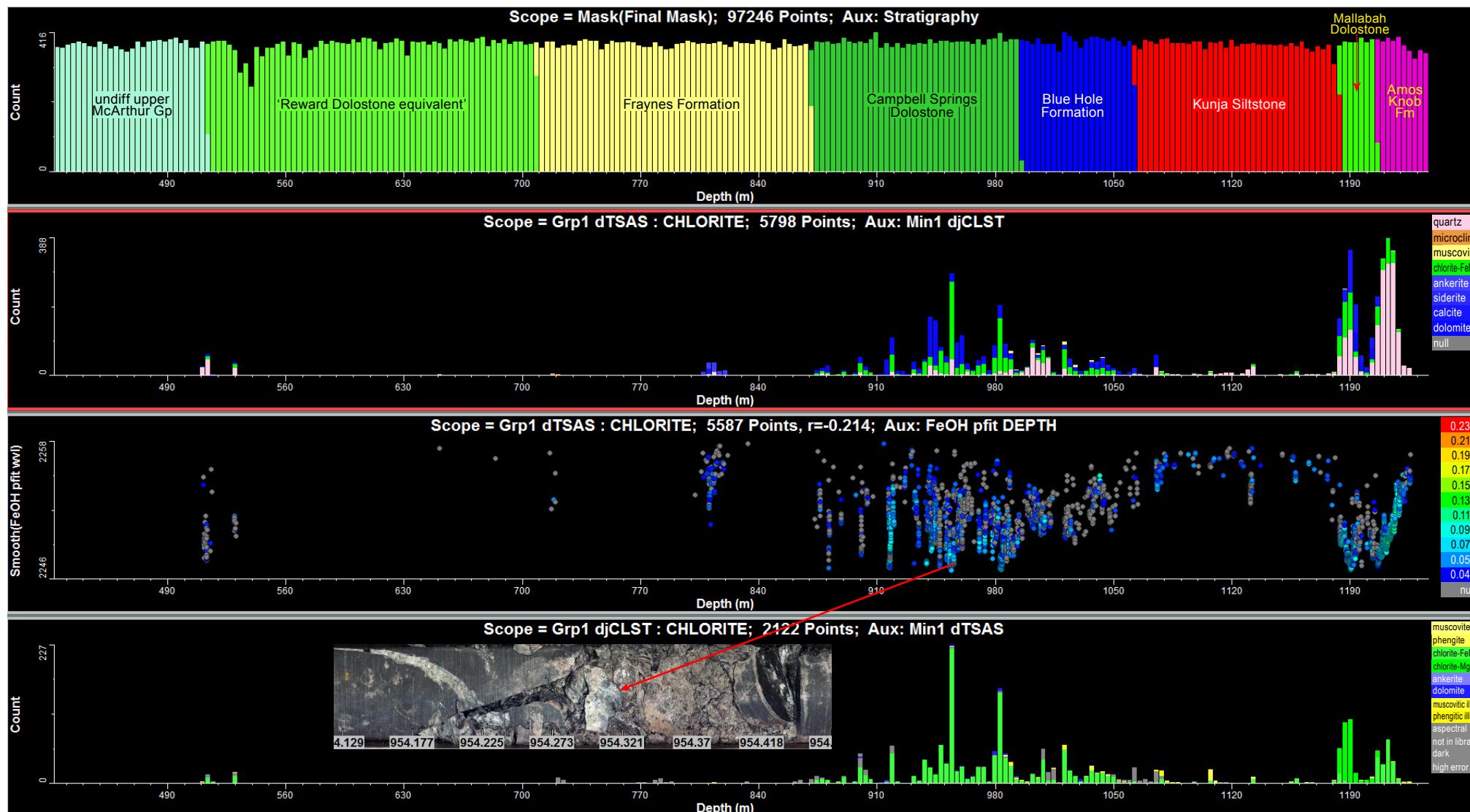


Row 1 shows the logged stratigraphic intervals from Pangaea (2015). Row 2 plots the SWIR spectra that match dominantly to white mica, coloured by the dominant TIR mineral. Row 3 are the SWIR white micas plotted by the wavelength of the white mica feature around 2200 nm and coloured by the depth of that feature. Wavelength changes may show white mica composition changes. The depth of the white mica feature is a measure of the strength (abundance) of the white mica. The lower part of the Fraynes Formation has an increased white mica abundance. White mica is also abundant in the Kunja Siltstone and Amos Knob Formation. Row 4 plots the SWIR spectra that match dominantly to white mica (masking out any spectra with <85% weighting of white mica), plotted by the white mica wavelength and coloured by the white mica crystallinity. Row 5 plots the TIR spectra that match dominantly to white mica, coloured by the dominant SWIR mineral. Some of the TIR white micas are aspectral in SWIR (grey colour, from around 855 m) and the core imagery (see inset image) indicates these are black shales (which are aspectral in SWIR due to dark / noisy spectra).



# Manbulloo S1: Chlorites

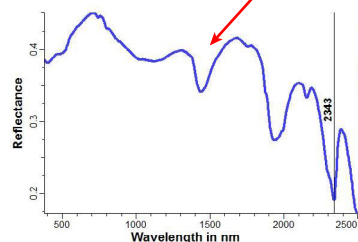
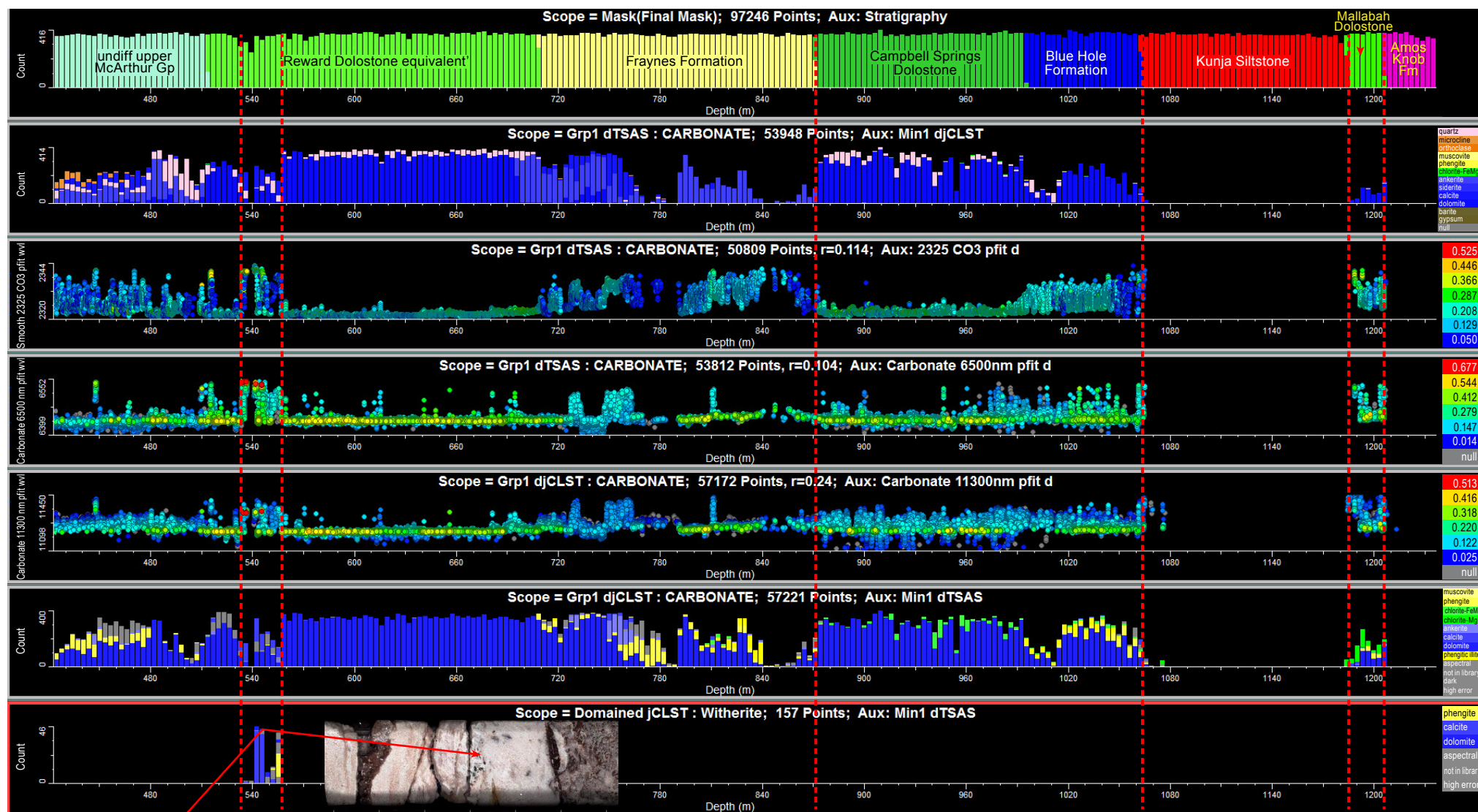
(View | Plot Layouts Load file 4\_VNIRSWIR: Chlorites)



Row 1 shows the logged stratigraphic intervals from Pangaea (2015). Row 2 plots the spectra that match dominantly to chlorite in the SWIR, coloured by the dominant TIR mineral. Chlorite in the Amos Knob Formation is commonly found with quartz. Row 3 plots SWIR chlorite spectra by the wavelength of the characteristic chlorite feature around 2255 nm and coloured by the depth of that feature. There is a range of chlorite composition, and the chlorite appears patchy. Row 4 plots the spectra that match dominantly to chlorite in the TIR, coloured by the dominant SWIR mineral. Chlorite is most common in the deeper part of Manbulloo S1. Chlorite in the Campbell Springs Dolostone is at specific depths (vertical dots). A closer look at one of these intervals (see inset image) indicates chlorite within a fault crush zone.

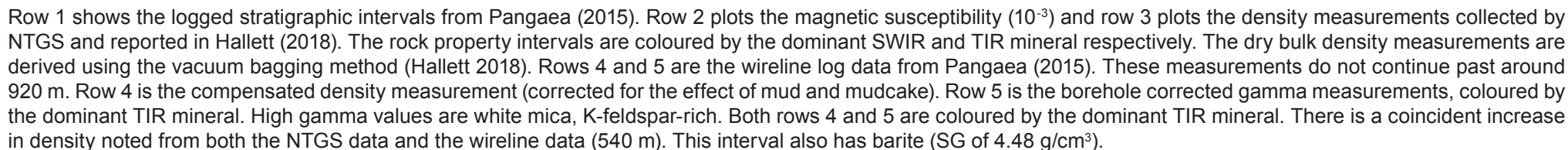
# Manbulloo S1: Carbonates

(View | Plot Layouts Load file 5\_VNIRSWIR: Carbonates)



Row 1 shows the logged stratigraphic intervals from Pangaea (2015). Row 2 plots the SWIR spectra that match dominantly to carbonate, coloured by the dominant TIR mineral. Row 3 are the SWIR carbonate spectra plotted by the wavelength of the characteristic carbonate reflectance feature around 2325 nm and coloured by the depth of that feature. Changes in wavelength are analogous to carbonate composition. Rows 4 and 5 plot TIR spectra that match dominantly to carbonate, plotted by the wavelengths of characteristic carbonate features in the TIR around 6500 nm and 11300 nm. The results are coloured by the strength (height) of that feature. Row 6 plots the TIR spectra that match dominantly to carbonate, coloured by the dominant SWIR mineral. Row 7 plots the TIR matches to witherite (Ba carbonate). Dotted lines highlight changes in carbonate abundance or composition. Note the zone with matches to witherite also has longer wavelength features in both the SWIR and TIR (rows 3 – 5; image inset left). The SWIR feature is around 2343 nm (calcite is 2340 nm). The witherite is a minor mineral component (calcite is dominant) in the TIR. Inset image (above) is of the barium-bearing carbonate.

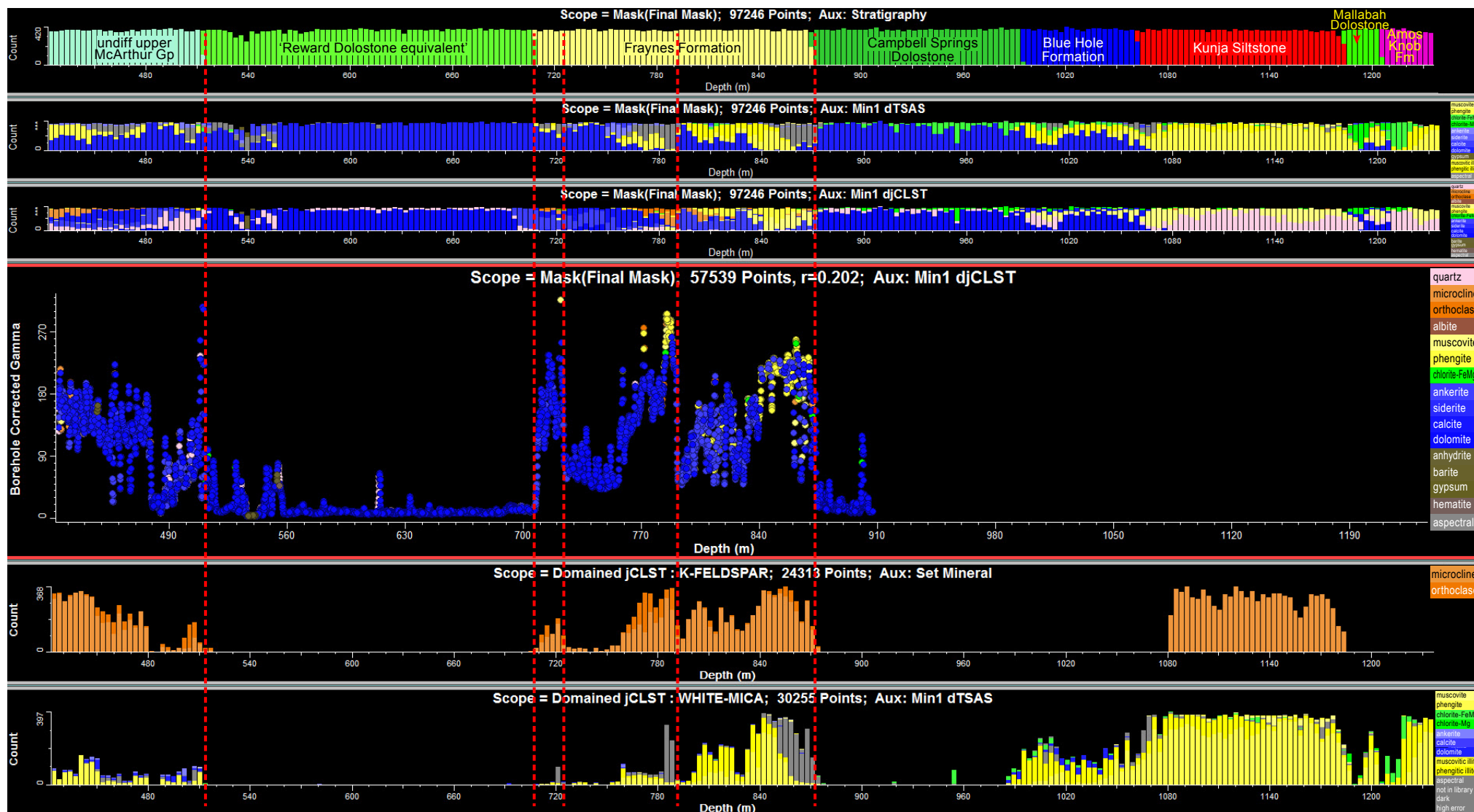
(View | Plot Layouts Load file  
6\_VNIRSWIR: Petrophysics)





# Manbulloo S1: Gamma Ray Values

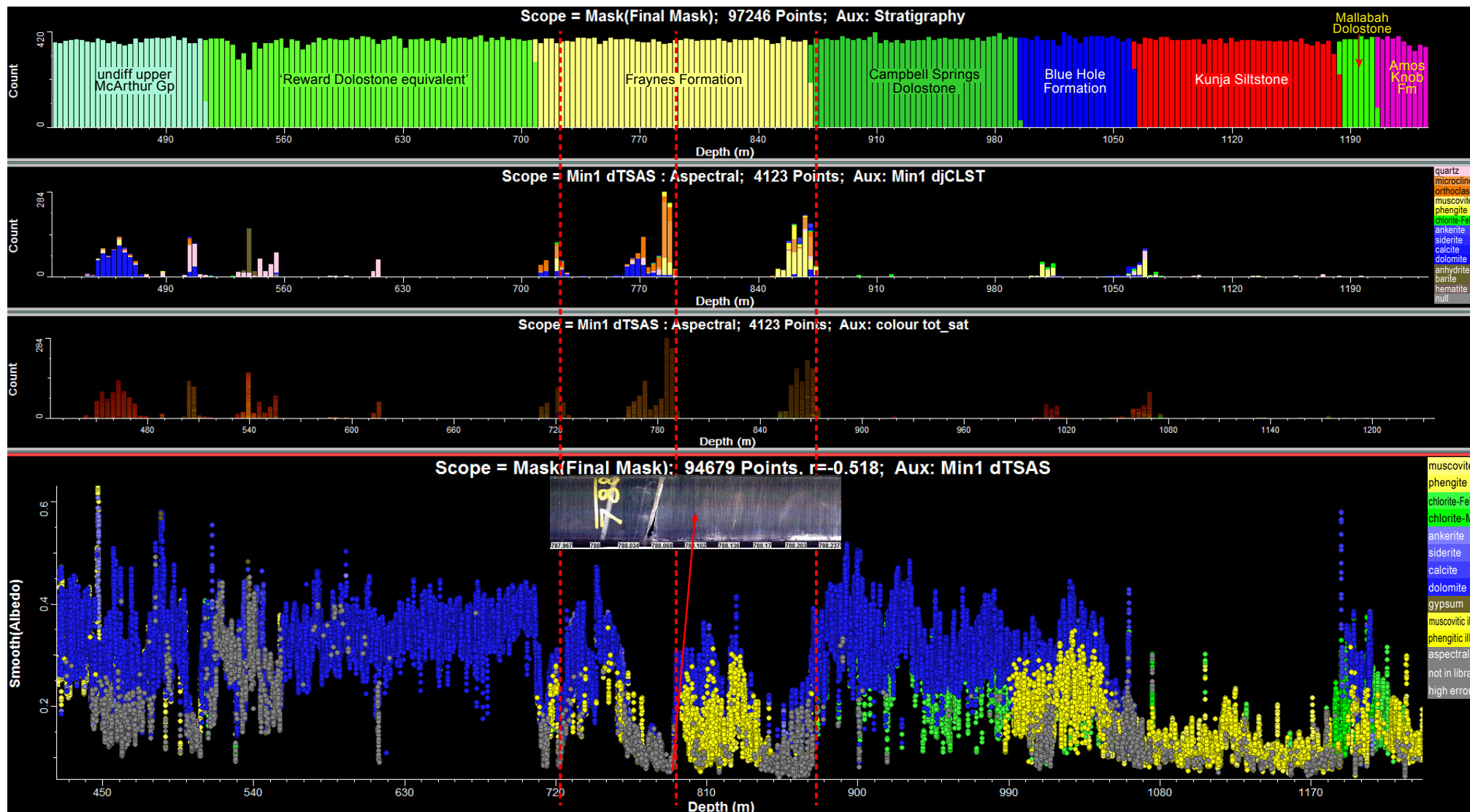
(View | Plot Layouts Load file 7\_VNIRSWIR: Gamma ray values)



Row 1 plots the logged stratigraphy from Pangaea (2015). Rows 2 and 3 plot the dominant mineral matches in the SWIR and TIR respectively. Row 4 plots the corrected borehole wireline gamma ray data from Pangaea (2015) and coloured by the dominant TIR mineral. The gamma ray data values had a top cut of 325 applied as there is one peak at 721 m with a max value of 855 that skews the data. The gamma values are not reported below 907 m. Note that the gamma peaks correspond to the stratigraphic boundaries to the base of the Fraynes Formation. Rows 5 and 6 plot the TIR spectra that have a match to K-feldspar and white mica respectively. As elevated gamma ray response can be from the presence of potassium, the K-feldspar and white mica matches are plotted below the gamma ray values. Dotted lines show areas with changes in both gamma ray response and K-feldspar / white mica presence. There appears to be a correlation between the gamma peaks and the presence of potassium-bearing minerals. Within the Fraynes Formation, there are 3 sharp peaks, with a decrease in gamma that corresponds to a decrease in K-feldspar / white mica content. The two deepest peaks in the Fraynes Formation are interpreted to be flooding surfaces; they may be analogous to the two flooding surfaces identified in the (stratigraphically equivalent) Barney Creek Formation (M.Kunzmann, pers comm).

# Manbulloo S1: Aspectral response in SWIR

(View | Plot Layouts Load file 8\_Aspectral in SWIR)



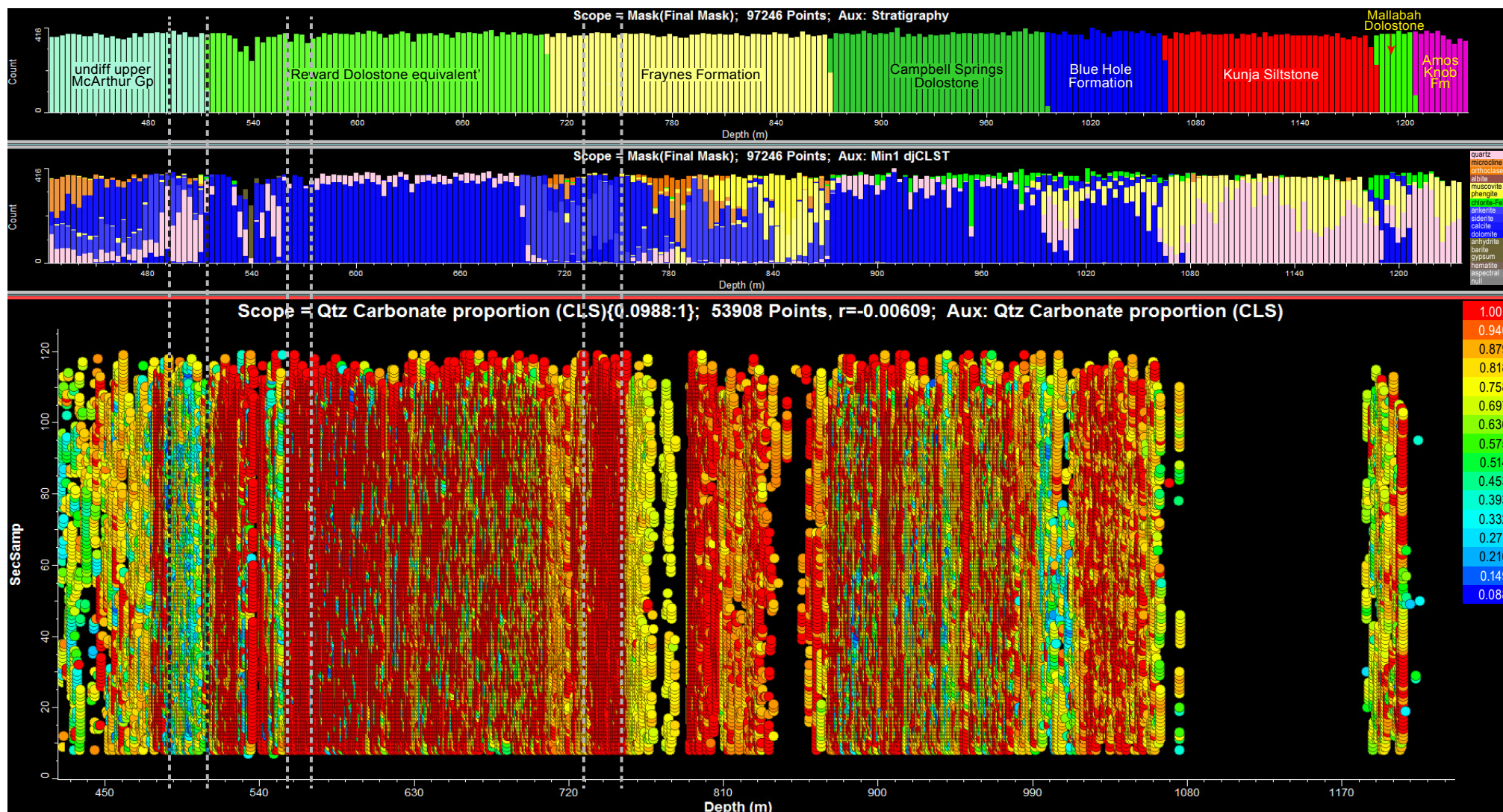
A SWIR aspectral response is when the SWIR spectra cannot match to the library mineral spectra. This may be due to noisy spectra (that may result from measurements on dark core), or measurements from core that contain minerals that are not in the TSA SWIR library. Some minerals do not have any diagnostic SWIR reflectance features, for example silicates such as quartz.

Row 1 shows the logged stratigraphic intervals from Pangaea (2015). Rows 2 and 3 plot the SWIR spectra that are classed as aspectral and coloured by the dominant TIR mineral (row 2) and the core colour (row 3). Note the highest number of SWIR aspectral occurrences in the Fraynes Formation commonly match to either K-feldspar or white mica in the TIR. Row 4 plots the smoothed albedo of all spectra (not just aspectral) and is coloured by the SWIR spectra (grey is aspectral). This plot is also on row 5, page 7. The dotted lines show where the SWIR aspectral response is highest in the Fraynes Formation. This highlights the black shales (grey dots with low albedo in Row 4). Some of the black shales (inset image) contain white mica (from the TIR) but the SWIR response is aspectral as the core is dark and so has a low reflectance (and higher noise) in these shorter wavelengths. These dotted lines are also at the same location as the gamma peaks and corresponding K-feldspar / white mica increases on rows 5 and 6, page 14.



# Manbulloo S1: Experimental carbonate-quartz proportions

(View | Plot Layouts Load file  
3\_TIR: QuartzCarbonateProportions)



Row 1 plots the logged stratigraphy from Pangaea (2015). Row 2 plots the dominant TIR mineral, using the domained jCLST. Row 3 is a carbonate-quartz proportion calculated from the CLS carbonate weights/[CLS carbonate weights + CLS quartz weights] (and masking out spectra that either do not have quartz and carbonate present or are 100% quartz, but allowing spectra that match 100% to carbonate). Row 4 plots 'sec samp' (section-relative sample number) as the y-axis so that each depth bin represents one tray row going from left to right (low SecSamp to high SecSamp) for each depth bin. Row 4 is coloured by the quartz-carbonate proportion with high values (red) being carbonate-rich and low values (blue) being quartz > carbonate. Blank areas are spectra that do not have both quartz and carbonate (eg may have carbonate and K-feldspar, or white micas, or chlorites). Black dotted lines in the 'undifferentiated upper McArthur Group' sediments show a quartz > carbonate interval. Grey dotted lines highlight high carbonate >> quartz zones in the 'Reward Dolostone equivalent' and the Fraynes Formation. Both of the highlighted intervals are below a sharp change in carbonate-quartz proportions. In the 'Reward Dolostone equivalent', there appears to be a sharp change to quartz-rich sediments. The imagery indicates a quartz-carbonate breccia interval. In the Fraynes Formation, the grey dotted carbonate-rich zone is immediately underlying a black shale at 723.6 m (with corresponding gamma peak, page 14). So the carbonate-quartz proportions can be used to indicate depositional changes in Manbulloo S1.



## Manbulloo S1: Summary of HyLogger data interpretation

- The stratigraphic picks for the upper stratigraphic units correlate with the borehole gamma peaks. These upper stratigraphic units do not have mineralogically distinct boundaries. The lower stratigraphic picks are mineralogically distinct. The exception is the boundary between the Campbell Springs Dolostone and the underlying Blue Hole Formation. This boundary is mineralogically gradational.
- Carbonate is the dominant mineral group, to 1063 m (base of the Blue Hole Formation). The Kunja Siltstone and the Amos Knob Formation are dominated by quartz and white mica, with no carbonate. This contrasts with the carbonate-bearing Kunja Siltstone and Amos Knob Formation in DD90VRB2 (Smith 2016). The Amos Knob Formation also contains carbonate in drillhole LBD1 (Smith 2015).
- Within the 'Reward Dolostone equivalent' (which may be the Killaloc Formation), a high density low magnetic susceptibility interval correlates with matches to TIR barite (which is spectral in the SWIR). Surrounding the barite matches are witherite / calcite (indicating high Ba content) and also ferroan dolomite (see image below).
- The Fraynes Formation may be stratigraphically equivalent to the Barney Creek Formation. Within Manbulloo S1, there are sharp gamma peak cycles that decrease moving uphole. These gamma peaks, which correlate with increased K-feldspar and white mica, are black shales. The lower two peaks in the Fraynes Formation may be analogous to the flooding surfaces identified in the Barney Creek Formation (M Kunzmann, pers comm).



Logged 'Reward Dolomite equivalent' with hematitic (ferroan dolomite?) on left, cut by a quartz vein running subparallel to core axis. The quartz vein is cross-cut by REE-bearing carbonate (mainly calcite?) from middle to right of image (downhole). Further downhole (~2 metres) are barite and barium carbonate (matches to witherite).

## References

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## Manbulloo S1: TSG metadata

File | Dataset Info

8446303\_Manbulloo\_S1\_tsg

Metadata | Sizes | Description | TSA Summary

Hole name: 8446303\_Manbulloo\_S1 | Logger: HyLogger 3-7

Project: National Virtual Core Library

Owner/Cust: Northern Territory Geological Survey

Author: Belinda Smith

Drilled: ☒ 2014-08-14 14:00 | Scanned: ☒ 2014-09-08 14:00

Latitude: 14.927675 | Long: 132.268419 | Datum: GDA94

Azimuth: 0.000000 | Incl: -90.000000 | RL: 144.800000

This dataset has a database entry.

Load From WFS

OK Cancel

8446303\_Manbulloo\_S1\_tsg

Metadata Sizes Description TSA Summary

Reprocessing to include trays missing from the original scan in 2014. Reinstated trays 34-40. Notes below from original processing: Checked spectra and turned off diaspore, saponite, topaz, Mg chlorite, epidote, zoisite, amphiboles, phlogopite, tourmalines, magnesite, alunites, jaorosite. Magnesite may be present; just not showing in TIR. Trace talc at 890.06m. After recal: turned off serpentine. Checked WNR and turned off artefacts. This version also updated to TSG8 and an extra domain for the included trays, which have significant barite, ferroan dolomite and Ni-bearing carbonate. Stratigraphic log from WCR in PR2015-0017. Added petrophysics and updated to TSG8.0.3.16 in January 2019.

OK Cancel

8446303\_Manbulloo\_S1\_tsg

Metadata Sizes Description **TSA Summary**

TSA set: ☒ SWIR ☐ VNIR ☐ TIR Copy to clipboard

Mineral	Sys %	Ustr %	Sys m	Ustr m
Dolomite	38.91	40.03	379.57	390.25
Phengite	17.91	17.87	173.67	173.14
Phengiticillite	5.72	5.65	55.20	54.63
Chlorite-Mg	4.82	0.90	46.27	8.65
Aspectral	3.13	3.35	31.13	33.32
Ankerite	2.58	1.98	25.16	19.40
Chlorite-FeMg	0.95	3.95	9.07	37.82
Calcite	0.93	1.06	9.10	10.37

OK Cancel

8446303\_Manbulloo\_S1\_tsg\_tir

Metadata Sizes Description TSA Summary

Turned off pyroxenes, olivines, garnets, plus SWIR mineral groups (dark micas, serpentine, amphiboles, epidotes, alunites, prehnite, pyrophyllite) as for SWIR. Updated to TSG8 with carbonate quartz proportion scalars in January 2019. Carbonate quartz proportion scalar calculated on CLS (due to mineral mixtures of up to 6 minerals) and masking out K-Feldspar, white mica and chlorite (to attempt to map the variations between quartz and carbonate content, not quartz with white mica, or carbonate with feldspar, for example. A high value (red colour, >1) would indicate almost all carbonate. Low values would indicate quartz >> carbonate. Experimental scalar only; use with caution.

OK Cancel

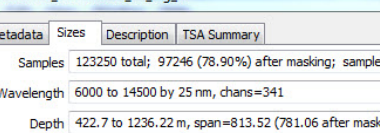
8446303\_Manbulloo\_S1\_tsg\_tir

Metadata Sizes Description **TSA Summary**

TSA set: ☐ SWIR ☐ VNIR ☒ TIR Copy to clipboard

Mineral	Sys %	Ustr %	Sys m	Ustr m
Dolomite	28.60	25.40	278.73	247.33
Quartz	16.25	16.82	158.16	163.66
Calcite	11.04	10.08	107.72	98.42
Muscovite	7.64	8.36	73.85	81.05
Microcline	4.42	5.12	43.00	49.81
Magnesite	2.37	0.00	23.06	0.00
Phenigite	2.27	2.75	21.95	26.56
Chlorite-FeMg	1.78	2.05	17.02	19.56
Orthoclase	1.27	0.70	12.39	6.92

OK Cancel



The screenshot shows a window titled '8446303\_Manbulloo\_S1\_tsg\_tir'. Inside, there is a table with four columns: 'Metadata', 'Sizes', 'Description', and 'TSA Summary'. The table contains the following data:

Metadata	Sizes	Description	TSA Summary
Samples	123250 total; 97246 (78.90%) after masking; sample=8mm		
Wavelength	6000 to 14500 by 25 nm, chans=341		
Depth	422.7 to 1236.22 m, span=813.52 (781.06 after masking)		
Scalars	System=34, core=13, user=61, total=108		
Linescan	Lines per sample=124, width=1170, JPEG quality=80		
Profilometer	Measurements per sample=128		
Disk size (MB)	Sp1:333, Sp2:511, img:1641, prof:60, pic:470, tot:3014.8		
Size / m (MB)	Sp1:0.41, Sp2:0.63, img:2.02, prof:0.07, pic:0.58, tot:3.6		

At the bottom of the window, there are 'OK' and 'Cancel' buttons.

From HyLogger Checklist icon

The Spectral Geologist (BELINDA SMITH) - 8446303\_Manbulloo\_S1.tsg

File Edit View Window Help

File Edit View Window Help [Icons: New, Open, Save, Print, Undo, Redo, Copy, Paste, Find, Zoom, Color, Style, Layer, View, Print, Save, Close, Quit]

HyLogging Checklist for 8446303\_Manbulloo\_S1\_tsg

Summary Basic TSA Scalars Domains & Plots DBase Journal

This dialog provides customers with the author's summary of steps taken and processing levels reached prior to their accessing this data. In addition, the Journal automatically tracks most changes and is worth visiting.

Signed off

Final mask: Yes	Depth logging: Yes
Imagery: Yes	System script scalars: Yes
Analyst's scalars: Yes	Imported scalars: Yes
User SWIR TSA: Yes	User TIR TSA: Yes
User VNIR TSA: Yes	Domains: Yes
Rockmarks: Yes	Plots: Yes
Layouts: Yes	Database: Yes

Save report

Include: ☒ Checklist ☒ Event journal ☒ as filtered

HyLogging Checklist for 8446303\_Manbulloo\_S1\_tsg

Summary Basic **TSA** Scalars Domains & Plots DBase Journal

User SWIR TSA

☒ Created ☒ Active minerals edited

☒ Through Domain RMS ☒ Noted in 'Dataset Info'

☐ Active minerals list exported ☒ Signed off by analyst

User TIR TSA

☒ Created ☒ Active minerals edited

☒ Through Domain RMS ☒ Noted in 'Dataset Info'

☐ Active minerals list exported ☒ Signed off by analyst

User VNIR TSA

☒ Created ☐ Active minerals edited

☒ Through Domain RMS ☒ Noted in 'Dataset Info'

☐ Active minerals list exported ☒ Signed off by analyst

OK Cancel

HyLogging Checklist for 8446303\_Manbulloo\_S1\_tsg

Summary Basic TSA Scalars Domains & Plots DBase Journal

**Imported Scalars**

☒ Geology ☐ Assays  
☒ Other ☒ Signed off by analyst

**Rockmarks**

Level of attention: ☒ None ☐ Moderate ☐ Significant  
☐ Validation XRD imported ☒ Signed off by analyst

**Standard Batch-script Scalars**

Interp support: ☐ Minor ☒ Major ☐ Essential  
☒ Through 'Final Mask' (or better) ☒ Assigned to appropriate groups  
☐ Ineffective scalars deleted ☒ Signed off by analyst

**Analyst's Scalars**

Interp support: ☐ Minor ☒ Major ☐ Essential  
☒ Through 'Final Mask' (or better) ☒ Assigned to appropriate groups  
☐ Ineffective scalars deleted ☐ Some aux-match scalars included  
☒ Signed off by analyst

OK Cancel

HyLogging Checklist for 8446303\_Manbulloo\_S1\_tsg

Summary Basic TSA Scalars Domains & Plots DBase Journal

**Domains**

- ☒ Created
- ☒ SWIR RMS lists hand-edited
- ☒ SWIR|TIR RMS lists aligned
- ☒ TIR CLS residual optimised
- ☒ Descriptions filled in
- ☒ TIR RMS lists hand-edited
- ☒ TIR CLS scalars created
- ☒ Signed off by analyst

**Plots**

- ☒ Reputable User / Domained TSA
- ☒ Significant imports
- ☒ All screens optimised
- ☐ Optimised TIR CLS
- ☒ Significant Batch / User scalars
- ☒ Signed off by analyst

**Layouts**

- ☒ Survey-standard layouts
- ☒ Each screen checked, all layouts
- ☒ Layout notes in Dataset Info
- ☒ Dataset-specific layouts
- ☐ Groundhog day for Viewer clients
- ☒ Signed off by analyst

OK Cancel



## HyLogger specifications

The TSG dataset originated from HyLogger™3–7. The HyLogger instrument rapidly measures reflectance spectra and also captures continuous high-resolution digital colour imagery of drill cores in their original trays.

HyLogger 3–7 was built by CSIRO (CSERE, North Ryde, NSW) and delivered to NTGS in February 2010 as part of the AuScope National Virtual Core Library (NVCL) project, a collaboration between Federal Government's Department of Innovation, Industry Science and Research, CSIRO and state and territory Geological Surveys.

The HyLogger has a continuous motion table that moves at 48 mm/second, and three spectrometers: a silicon-detector grating spectrometer for the [380, 1072] nm VNIR interval, an InSB-detector FTIR (fourier transform infrared) spectrometer for the [1072, 2500] nm SWIR interval and a further FTIR spectrometer with a HgCdTe photoconductive detector for the [6000, 14500] nm TIR interval. The spectrometers measure 12 spectra per second, or one spectrum for each 4 mm at the standard table speed of 48 mm/second. The camera, a Basler piA1900-32gc, takes 12 frames per second (or one for every 4 mm).

Full details of the HyLogger specifications can be found in Schodlok *et al* (2016b).

# Glossary

Glossary of acronyms and technical terms commonly used in HyLogging spectroscopy.

albedo	Normally applied to the mean broadband brightness of a spectrum over a specified wavelength range. A white or altered sample will commonly have a high albedo, whereas a graphitic rock will have a very low albedo.
aspectral	An aspectral response is a spectrum that does not match a TSA library spectrum within the SRSS error cut-off. An aspectral response may be due to many different factors including: dark/noisy spectrum; a mineral not in the TSA library; a silicate mineral without any absorptions in the SWIR (such as olivines, pyroxenes, feldspars, quartz without fluid inclusions).
AlOH	Aluminium hydroxide.
AusGIN	Australian Geoscience Information Network (Geoscience portal): <a href="http://portal.geoscience.gov.au/gmap.html">http://portal.geoscience.gov.au/gmap.html</a> is a web portal that hosts NVCL data.
AuScope	The national provider of integrated research infrastructure, of which the NVCL is an infrastructure programme. The AuScope portal ( <a href="http://portal.auscope.org/portal/gmap.html">http://portal.auscope.org/portal/gmap.html</a> ) hosts NVCL data.
CLS	Constrained Least Squares – an alternative unmixing classifier that uses a RMS to minimise non-unique mineral modelling. Used mainly to model TIR spectra that can have several mixed mineral matches.
Corstruth	A webpage that plots the results (as a pdf summary) from an automated analysis of HyLogger data in the NVCL; <a href="http://www.corstruth.com.au">www.corstruth.com.au</a>
domain	A zone within a drillhole interpreted to contain a restricted set of minerals that are different to adjacent zones. Unmixing algorithms applied to domained datasets use a RMS that has been defined for each domain by the processing geologist.
FTIR	Fourier transform infrared spectrometer.
HgCdTe	Mercury cadmium telluride used in infrared detectors.
HQ	Hull quotient – a type of background corrected spectrum.
InSb	Indium antimonide – used in infrared detectors.
MCT	Mercury cadmium telluride used in infrared detectors.
MgOH	Magnesium hydroxide.
nm	Nanometre, being one billionth of a metre. A HyLogger 3 operates between 380 and 14 500 nm, with no measurements between 2500 to 6000 nm.
NVCL	National Virtual Core Library; the library of nationally available TSG datasets
scalar	Any set of imported or calculated values associated with spectral data loaded in TSG.
RMS	Restricted mineral set. The processor limits the set of possible mineral matches based on the geological understanding and spectral characteristics of the domain.
SEM	Scanning Electron Microscopy is a type of electron microscope that images the sample surface by scanning it with a high energy beam of electrons, giving information on sample composition and other properties. SEM results may be used to validate mineral identification by the HyLogger.
SNR	Signal-to-noise ratio.
SRSS	Standardised residual sum of squares (TSA's measure of mineral identification error). Low SRSS values are more reliable than high ones. The current 'bad' threshold is 1000.
SWIR	Shortwave infrared (light). In HyLogging applications it nominally covers the range 1000–2500 nm.
TSA	'The Spectral Assistant' – CSIRO trademarked algorithm that uses training libraries of pure spectra to match an unknown spectrum to a single mineral or to identify mixtures of two or more minerals. Part of the TSG software package.
TSG	'The Spectral Geologist' – CSIRO-developed specialist processing software, designed for analysis of field or laboratory spectrometer data. <a href="http://thespectralgeologist.com/">http://thespectralgeologist.com/</a>
TIR	Thermal infrared (light). In HyLogging applications it nominally covers the range 6000–14000 nm.
VIS	Visible (light). The human eye is nominally sensitive between 390 and 750 nm.
VNIR	Visible near infrared (light). In HyLogging applications it nominally covers the range 380–1000 nm.
volume scattering	Radiation that is reflected after some absorption into the rock and changes the spectral shape and features. TIR spectral interpretation assumes that there is only surface scattering in a spectrum. Volume scattering leads to errors in TSA and CLS modelling.
wvl	Wavelength - used in TSG scalar names.
XRD	X-ray diffraction - an analytical technique that reveals information about the crystallographic structure, physical properties and chemical composition of a sample. It is based on observing the scattered intensity of an X-ray beam hitting a sample and measuring the scattered angle and wavelength or energy.

# Guide to scalars in figures produced using TSG software

The terms used in the titles, x and y-axis for figures produced from TSG are described in the table below:

2200 wvl 2nd deepest	FEATEX scalar that measures the wavelength of the second deepest absorption feature from 2200nm +/- 50nm. Designed to measure the wavelength of the kandite doublet, which has a variable wavelength depending on whether it is kaolinite or dickite.
2325 CO3 PFIT wvl	PFIT scalar to measure the wavelength of a trough minima between 2290–2370 nm with a depth >0.05; polynomial order 8; hull envelope divided by reflectance reported as wavelength at minimum in nm. Used mainly to analyse carbonate composition changes by observing wavelength changes in the dominant absorption feature for carbonate in the SWIR.
Al smectite abundance	Developed by CSIRO in 2011 as multiple feature extraction method (MFEM) batch script, this (unvalidated) scalar maps montmorillonite and beidellite abundance by measuring the continuum removed depth of a fitted 4th order polynomial between 2120 and 2245 nm.
Apatite 9200 PFIT d	PFIT scalar created by J Huntington to confirm the TSA apatite response. Measures the wavelength of the minimum trough between 9192 nm and 9270 nm with a depth of >0.0006; polynomial order 6; hull envelope subtract base reflectance to give a relative depth.
Aux	Aux in a plot indicates the parameter that is colouring the points (bars in bar plot, points in scatter plot) in a figure. For example, Aux: stratigraphy indicates that the colours relate to stratigraphy. The key to the Aux colours are on the right side of each plot.
Aux match scalar	Aux matching involves simple curve matching between spectra in a main dataset and spectra in a Aux (Auxiliary or Custom) dataset. The Aux dataset is usually a custom library containing special hand-chosen spectra that have been interpreted in detail.
Carbonate 6500 nm wvl	Experimental batch scalar created by CSIRO derived from the reflectance of the 6500 nm wavelength peak. Used to determine differences in the wavelength of the peak around 6500 nm, which shifts with different carbonate compositions.
Carbonate 11300 nm wvl	PFIT scalar to measure the wavelength of the peak maxima between 11000–11580 nm with a height of >0.04; polynomial order 9; hull envelope subtract base reflectance to give wavelength at maximum.
Christiansen Minimum	Experimental batch scalar created by CSIRO that plots the Christiansen Minimum wavelength. The Christiansen Minimum occurs when the refractive index of the sample approaches the refractive index of the (medium) air surrounding the mineral grains, resulting in minimal scattering and minimal reflectance (Conel 1969). The Christiansen Minimum wavelength varies according to composition, so measuring the Christiansen Minimum wavelength can differentiate igneous rock compositions in the TIR.
Colour tot_sat	TSG standard scalar; it calculates the colour (separately per band) from the visible interval of the reflectance spectra and it is enhanced by a 'total saturation' (the S band is 'wired to 1'; no pastels). Refer to TSG Help Manual for more explanation.
Count	The feature frequency plots are bar plots with y-axis = count. The count is cumulative number of features within a bin. The bin size will vary according to the x-axis, which might be depth, wavelength in nanometres etc.
FEATEX scalar	FEATEX scalars use a feature extraction algorithm in TSG to calculate the depth, width and/or wavelength position of a spectrum's absorption features. It uses pre-calculated feature extraction information from TSG's default algorithm.
Felsic-Mafic Index wvl	Experimental batch scalar created by CSIRO that maps the peak wavelength between 7500 and 12000 nm from a 4th order polynomial. Shorter wavelengths are more felsic than longer mafic ones. Most carbonate-bearing samples are excluded.
FeOH PFIT depth	PFIT scalar to measure the depth of a trough minima between 2240–2270 nm with a depth >0.04; polynomial order 10; hull envelope divided by reflectance reported as relative depth.
FeOH PFIT wvl	PFIT scalar to measure the wavelength of a trough minima between 2245–2260 nm with a depth >0.04; polynomial order 10; hull envelope divided by reflectance reported as wavelength at minimum in nm. Used mainly to analyse chlorite composition changes.
FeOx intensity (alt)	TSG standard scalar (batch scalar) that ratios the reflectance at 742 nm / reflectance at 500 nm to give the Fe slope.
Garnet 11400 comp wvl	PFIT scalar to measure the wavelength of the trough minima focussed between 10550 to 11600nm with a depth >0.2; polynomial order 6; hull envelope divided by base reflectance reported as wavelength at minimum in nm. Used to validate garnet species; almandine has shorter wvl; andradite has longer wvl.
Grp 1 Min	Group (coarse level) index of the primary mineral group component in a TSA result. Mineral groups include carbonates, white micas, pyroxenes etc. Grp2 Min would be the secondary/minor mineral group component in a mineral mix.

(continued on next page)



## Guide to scalars in figures produced using TSG software (continued)

Hydrocarbon presence	A PFIT scalar designed to measure the presence of an absorption feature at 1730 nm, which is often found associated with oil bleeds. Another scalar designed to identify hydrocarbons is the 23140 nm PFIT scalar. Using both scalars together can identify oil bleeds when the spectral response is preserved (it can deteriorate over time).
Kaolin composition	A CSIRO-built batch scalar in TSG that measures the composition and crystallinity of kaolin group minerals ranging from well-ordered kaolinite to halloysite to dickite (and nacrite).
Kaolinite PFIT 2160 doublet d	A PFIT scalar measuring the relative depth of the kandite doublet absorption feature, as an analogue for crystallinity / abundance of kaolinite. Used to confirm that TSAS-assigned well-crystalline and poorly-crystalline kaolinite are present. PFIT used hull envelope divided by reflectance; focussed on 2155–2180 nm to determine the relative depth using a >0.01 cut-off, polynomial order 3, masked through Final Mask.
Mask (Final Mask)	Mask scalars are used to filter out unwanted spectra caused by scanning tray edges, core blocks etc. TSG uses the Final Mask as the default mask for both SWIR and TIR datasets and will synchronise the mask for both datasets. Many in-built TSG scalars are calculated after being filtered through the Final Mask.
Min 1	Mineral index of the primary mineral for a TSA singleton match or primary mixture component. Min 2 is the subordinate/minor mineral in a TSA mineral mix.
PFIT scalar	PFIT scalars take a section of the spectrum specified by the user, optionally does a local continuum removal, fits a polynomial and calculates a result directly from the polynomial's coefficients. PFIT scalars are used to define the wavelength of noted spectral features.
Prehnite 1475 nm PFIT	PFIT scalar to measure the wavelength of a trough minima between 1460–1485 nm with a depth >0.009; polynomial order 6; hull envelope divided by reflectance reported as a relative depth. Used to confirm prehnite, with the depth of the 1475 nm feature indicative of abundance.
Quartz 8625 PFIT d abundance	Experimental PFIT scalar to measure the 'abundance' of quartz in a sample by measuring the depth of reflectance minima at 8625 nm, which is characteristic of the presence of quartz. Scalar measure returns relative depth in nm, by subtracting the low side of the minima from normalised TC reflectance using a depth >0.02 between 8580–8700 nm.
Quartz 8625 PFIT d MAV	Smooths the 8625 PFIT d abundance scalar (above) by using the mean through a moving window. The output smooths out the effect of outliers to display gross changes in the quartz abundance in plots.
Quartz absorption depth	Experimental batch scalar created by CSIRO to measure the depth of the characteristic quartz reflectance feature at 8625 nm. Similar to the Quartz 8625 PFIT d abundance scalar, but can be more effective in masking out spurious matches to some sulphates that formed from the core decomposition after drilling (refer Sever No.1 drillhole).
Quartz_H2O	An inbuilt batch scalar found in HotCore. Described as 'normalised ratio that maps samples with appreciable (1950 nm) water absorption in fluid inclusions, found mostly in quartz (and some carbonates)'.
Scope	The Scope option allows users to filter their data to visualise the behaviour of selected classes (eg; stratigraphy, mineral groups) and samples in different XY plots. The Scope indicates how many samples out of the total samples in the dataset are currently displayed in this plot window.
Smooth (Albedo)	TSG standard scalar (batch scalar) that first calculates the reflectance albedo over 450–2450 nm with basic channel outlier masking, then averages the numeric response (smooths) of the albedo. May also be called Albedo Rmean Smooth or Smoothed Albedo.
Smoothed scalar	Created by 'smooth an existing scalar using a moving window'. Generally uses averaging of the numeric response to create a smoothed scalar.
sTSAS, uTSAS, uTSA+, dTSA+	Mineral result from matching to the short wave infrared (SWIR) spectra against the TSA library. In TSG versions 7 and earlier; 'sTSAS' is the default system match. 'uTSAS' is the author-derived result from manually excluding some minerals and artefacts (eg; wooden core blocks, plastic chip tray spectra) during processing. In TSG versions 8 and later, TSA+ uses some information derived from selected features in both the SWIR and TIR to make a more informed choice about mineral mixtures. The 'd' indicates the results are 'domained'.
sTSAT, uTSAT, dTSAT	Mineral results from matching to the thermal infrared (TIR) spectra against the TSA library. 'sTSAT' is the default system match. 'uTSAT' is the author-derived result from manually excluding some minerals and artefacts during processing. 'dTSAT' indicates the results are 'domained'.
jCLST, ujCLST, djCLST	The jCLST algorithm has been developed by Andy Green (OTBC Pty Ltd; <a href="http://www.corstruth.com.au">www.corstruth.com.au</a> ) as a replacement for sTSAT, which unmixes the TIR spectra on a sample by sample basis without reference to the results in the VNIR or SWIR (which can commonly return spurious mineral matches). In comparison, jCLST interprets TIR data using the results from a modified TSAT, TSA+ and from scalars using selected features in the VNIR and TIR. 'ujCLST' is the author-derived results from manually excluding some minerals during processing. 'djCLST' is author-derived results from manually domaining the drillhole into zones of similar mineralogy and restricting the minerals available for the jCLST algorithm in each domain.

(continued on next page)

## Guide to scalars in figures produced using TSG software (continued)

sTSAV, uTSAV	Mineral result from matching to the visible near infrared (VNIR) spectra against the TSA library. 'sTSAV' is the default system match. 'uTSAV' is the author-derived result from manually excluding some minerals and artefacts (eg; galvanised tray spectral matches) during processing.
TIR-CLS1_CLS_min_1	CLS scalar showing the dominant modelled mineral (using the CLS unmixing algorithm) from the TIR wavelength range. For this scalar, the number of minerals allowed in the CLS mineral output is 3 (shows the 3 most dominant) although the scalar can allow for up to 6 minerals. The minerals available for modelling in the domain (RMS) is selected during the interpretation / processing stage.
TIRDeltaTemp	An inbuilt TSG scalar that measures the change in temperature between the instrument response measured from the rock and the background response. Plotting this scalar can highlight sulphides or artefacts (such as metal tray edges, metal depth marker tabs or instrument issues).
Tourmaline PFIT 2366 nm	PFIT scalar to measure the wavelength of a trough minima between 2360–2375 nm with a depth >0.015; polynomial order 5; root mean square error (RMSE) ≤ 0.06; hull envelope divided by reflectance reported as a relative depth. Used to search for tourmaline in mixtures with chlorite and white micas (which may have overlapping features at around 2206 nm and 2244 nm).
uTSA*	The result from TSA. The prefix 'u' is for 'user' and refers to the fact that TSA is trained on a reference library of minerals that have been limited by the author from the system set of minerals called sTSA* during the processing of the dataset. The minerals that are excluded from being matched to the TSA library are those that are considered to be unlikely in that geological environment and do not visually match the spectra well.
uTSAT invalid	A scalar created to mask out both Final Mask and 'aspectral', 'noisy' or 'null' for uTSAT Min 1 minerals. Plots that use uTSAT plot only spectra that were successfully matched in the TSA library, so noisy spectra (which may be noisy due to rubbly core, volume scattering etc) don't detract from displaying the dominant mineral or mineral group in the TIR. If a hole has a lot of 'invalid' spectra due to rubbly core, it may over-emphasise minor minerals in the TIR that are perhaps within unbroken core and may not reflect accurate mineral proportions.
White mica PFIT wvl	PFIT scalar to measure the wavelength of a trough minima between 2190–2229 nm with a depth >0.04; polynomial order 10; hull envelope divided by reflectance reported as wavelength at minimum in nm. Used mainly to analyse white mica composition changes by observing wavelength changes in the dominant absorption feature for white mica in the SWIR.
White mica PFIT d	PFIT scalar to measure the depth of a trough minima between 2190–2229 nm with a depth >0.12; polynomial order 10; hull envelope divided by reflectance reported as relative depth.
WM crystallinity	Arithmetic scalar measuring D2200 white mica divided by D1900 masked by the Final Mask. White mica crystallinity measures the depth of the AIOH absorption feature relative to the depth of the water feature. A deeper water feature indicates lower crystallinity and may indicate an illitic white mica (which may also have some compositional substitution).

### 1.1 Basic HyLogging Product Levels

0. **Machine Data** package (QCed & archived by collecting team / agency: i.e. all repeats taken care of and data checked).
1. **TSG Data** package - TSG imported and formatted data (see note about TSG-QC outputs)
  - 1A. TSG imported imagery, spectra and supporting data (nothing else done).  
Raw system TSA run on import but no checking. Depths only based on tray starts & ends.
  - 1B. Final masked, basic depth-logged data, imagery enhanced, new tray imagery & mosaics created.. Further updates possible.
  - 1C. All standard "system" scalars (includes basic masked and reviewed TSA mineralogy) created & checked.
  - 1D. User TSA results included (i.e. retrained TSA) and all Scatter screens changed to uTSAS.  
Minimum database entry point.
  - 1E. Non-standard mineralogical (manually-generated) "user" scalars added, thresholded and checked. Might include an Aux match library or stats (PC) analysis.
  - 1F. All metadata tables updated. Optimum database loadable level. Further updates possible.
2. **Integrated Data** package - Imported numeric or class scalars added into TSG and depths adjusted if required to fit assay intervals.
3. **Published Data** package - Signed-off for public (NVCL) database publication. Default set of products (for web discovery) identified and tagged.
4. **Down-sampled Data** package - Optional down-sampled version of all of above.
5. **Project data** package. Abstracted data from many holes integrated in some way.

The HyLogger Product level refers to the level of processing of a dataset. This dataset is at 'Level 3' as it has imported stratigraphic information.