



MICAP-OES 1000

Multi-elemental high throughput analysis of in-service coolants with a novel N₂ ICP-OES system following ASTM Method D6130

Introduction

Monitoring engine coolants has recently become a very important tool in tracking the health of the critical liquid coolant process in gas and diesel engines. The elemental composition of the coolant fluids provides important information regarding the corrosion protection and stability of these essential engine fluids. Maintaining effective cooling throughout an engine directly impacts the usable lifetime of these engines.

ICP-OES has been used routinely in the monitoring of in-service engine oils for many years, and recently has been applied to the monitoring of engine coolants. This note describes the use of the simultaneous MICAP N2 ICP-OES system for routine analysis of glycol-based coolants following the ASTM Method D6130, which is designed to monitor both new and in-service coolants by ICP-OES. For this work, high throughput sample introduction automation is used to optimize and speed up both the uptake and washout of these critical fluids for an easy multi-elemental analysis of the additive, corrosion, and contaminant elements.

Instrumentation

As engine coolants are aqueous-based liquids, the Radom Instruments MICAP-OES 1000 was configured with an aqueous sample introduction system. A double-pass cyclonic spray chamber and high-solids concentric nebulizer were coupled to the standard one-piece torch to introduce the samples into the N₂ plasma.

Microwave energy in the MICAP is coupled into the Cerawave™ ring in a highly efficient process that creates the magnetic fields required to inductively couple energy into the robust N₂ plasma emission source. The 4 MP sCMOS camera simultaneously collects the emission lines from the high-resolution spectrometer to detect the trace elemental signals.

The composition of these coolants can vary greatly based on their use, which alters physical properties such as viscosity and surface tension. To ensure consistent uptake and rinse-out of these coolant samples, an Elemental Scientific (ESI, Omaha, NE, USA) 4DXCi autosampler and SampleSense FAST discrete sampling valve was coupled to the MICAP.

This automated sampling valve quickly vacuum loads the sample into a loop while the valve optically monitors the arrival of the sample. Once the sample loading is complete (approx. 3 sec), the analysis is automatically triggered to start the collection of the emission spectrum from the plasma. Upon analysis completion, the next sample is automatically loaded while the nebulizer and spray chamber are rinsed out in preparation for the next sample analysis. This automated sampling valve provides the following advantages to the analysis:

- Sample uptake time is greatly shortened
- Uptake of each sample is optimized as coolants of varying viscosities are each optimally loaded
- Washout of coolants is enhanced as the spray chamber rinse is performed simultaneously along with the next sample uptake

Experimental Conditions

The analyses were performed on the MICAP-OES 1000 simultaneous N₂ ICP-OES system utilizing the conditions listed in Table 1. The settings used with the ESI SampleSense FAST automation system are also provided here. A complete view of this integrated analysis solution is displayed in Figure 1.

The analysis conditions employed on the MICAP follow closely the procedures outlined in ASTM Method D6130-24, Determination of Silicon and Other Elements in Engine Coolant by ICP-AES¹. Table 2 provides specifics on the analyte and internal standard (Co) wavelengths used for this analysis.

Table 1. MICAP conditions and FAST automation settings

Parameter	Value
Torch	Quartz 1-piece, 1.5mm injector
Spray Chamber	Double pass cyclonic
Nebulizer	High solids concentric glass
Sample Tubing	Blk/Blk PVC (0.76 mm ID)
Drain Tubing	Blu/Yel PVC (1.52 mm ID)
SampleSense FAST Loop	1 mL
FAST Carrier / Rinse	DI Water
Coolant Gas Flow	14 L/min
Auxiliary Gas Flow	0.4 L/min
Nebulizer Gas Flow	0.7 L/min
Peristaltic Pump	30 rpm
Plasma Viewing	Axial
Camera Exposure	10 sec (200 ms @ 50 reps)
# of Repeats	3

Table 2. Analyte and internal standard wavelengths

Analyte Wavelength (nm)	Internal Standard Wavelength (nm)
Al 396.152	Co I 240.725
B 249.677	
Ca 396.847	Co II 238.892
Cu 324.754	Co I 240.725
Fe 259.940	Co II 238.892
K 404.414	Co I 240.725
K 766.490	Co II 238.892
Mg 280.270	
Mo 317.034	Co I 240.725
Na 568.819	
P 213.618	Co II 238.892
Pb 283.305	Co I 240.725
Si 251.611	
Sn 283.998	
Zn 213.857	Co II 238.892



Figure 1. MICAP, SampleSense FAST valve & 4DXCi autosampler

Table 3. Analytes and calibration standards, in mg/L (ppm)

Element	Std 1	Std 2	Std 3	Std 4	Std 5	Std 6	CCV1	CCV2	CCV3
Al	0.2	2	20				2		
B	0.2	5	100					50	
Ca	0.2	2	20				2		
Cu	0.2	2	20				2		
Fe	0.2	2	20				2		
K	0.5	20	500				5	200	
Mg	0.2	2	20				2		
Na	20	200	1000					200	
P	5	100	200					100	
Pb	0.5	2	20				5		
Zn	0.2	2	20				2		
Mo				0.2	5	100			5
Si				0.2	2	20			2
Sn				0.5	2	20			5

Standard and Sample Preparation

The calibration standards, blanks, and samples were prepared in accordance with the protocols outlined in ASTM Method D6130-24¹. Multielement standards were prepared from aqueous standards (Inorganic Ventures, Christiansburg, VA, USA). Final standards and blanks were prepared in 18 MΩ deionized water (DI) and 5% ethylene glycol w/v (Sigma-Aldrich, St. Louis, MO, USA). See Table 3 above for the specific concentrations utilized. Note that since K was cross calibrated using two emission lines, a mid-point calibration check was performed for each line to confirm stable response during the analysis.

Thirty in-service coolant samples were sourced from a local heavy equipment supplier laboratory. These were augmented with two new coolant samples sourced commercially. Coolant supplied at usage levels were prepared by diluting 1:10 fold (v/v) in DI water, while concentrated (new) coolant samples were diluted 1:20 fold.

All standards and samples included the element Cobalt (Co) at 10 mg/L (ppm) as an internal standard (Inorganic Ventures). Additionally, 1% (w/v) Cesium (as CsCl, Sigma-Aldrich) was also added to all solutions. The Cs serves as an ionization buffer to eliminate matrix interference observed for coolant samples containing very high Na and K levels. Both the Co and the Cs were added into the diluent to facilitate the sample and standard preparation.

Results

The 30 in-service and 2 new coolant samples were successfully analyzed with MICAP and the results compiled. Figure 2 demonstrates the differences in the measured analytes across 2 in-service and one new extended lifetime coolant (ELC).

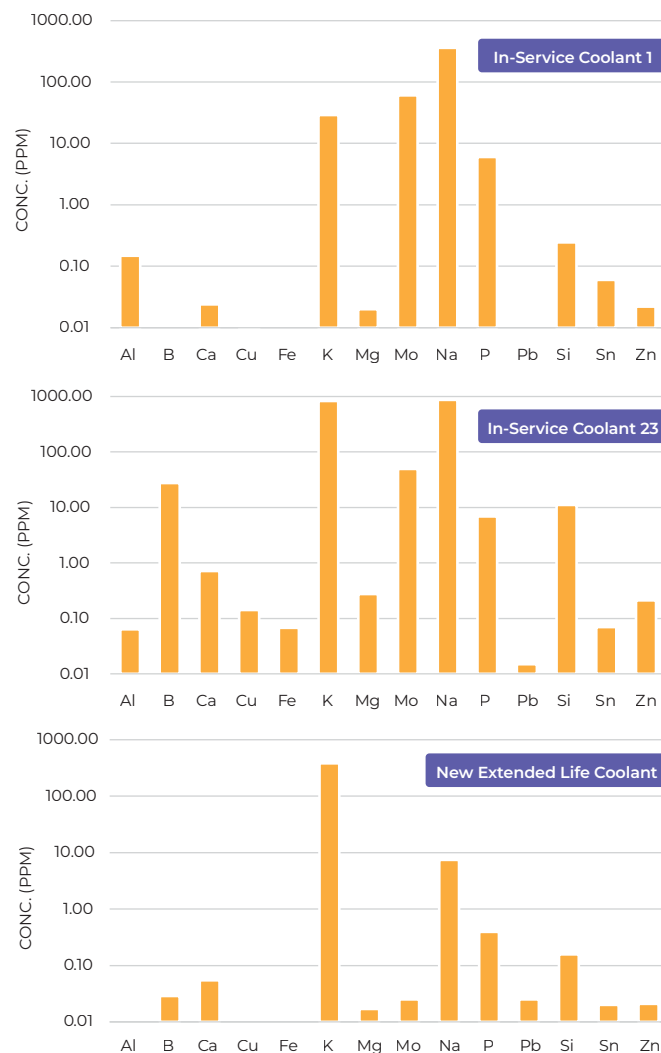


Figure 2. Comparison of elemental concentrations across three coolant samples analyzed

These coolants vary significantly in their makeup. They differ in color from various shades of red, orange, and green (see Figure 3). From general observation, the viscosities also vary from coolant to coolant. It was additionally

interesting to observe the wide range of concentrations of elements in various coolant samples. Figure 4 displays the K concentrations across the 32 samples.

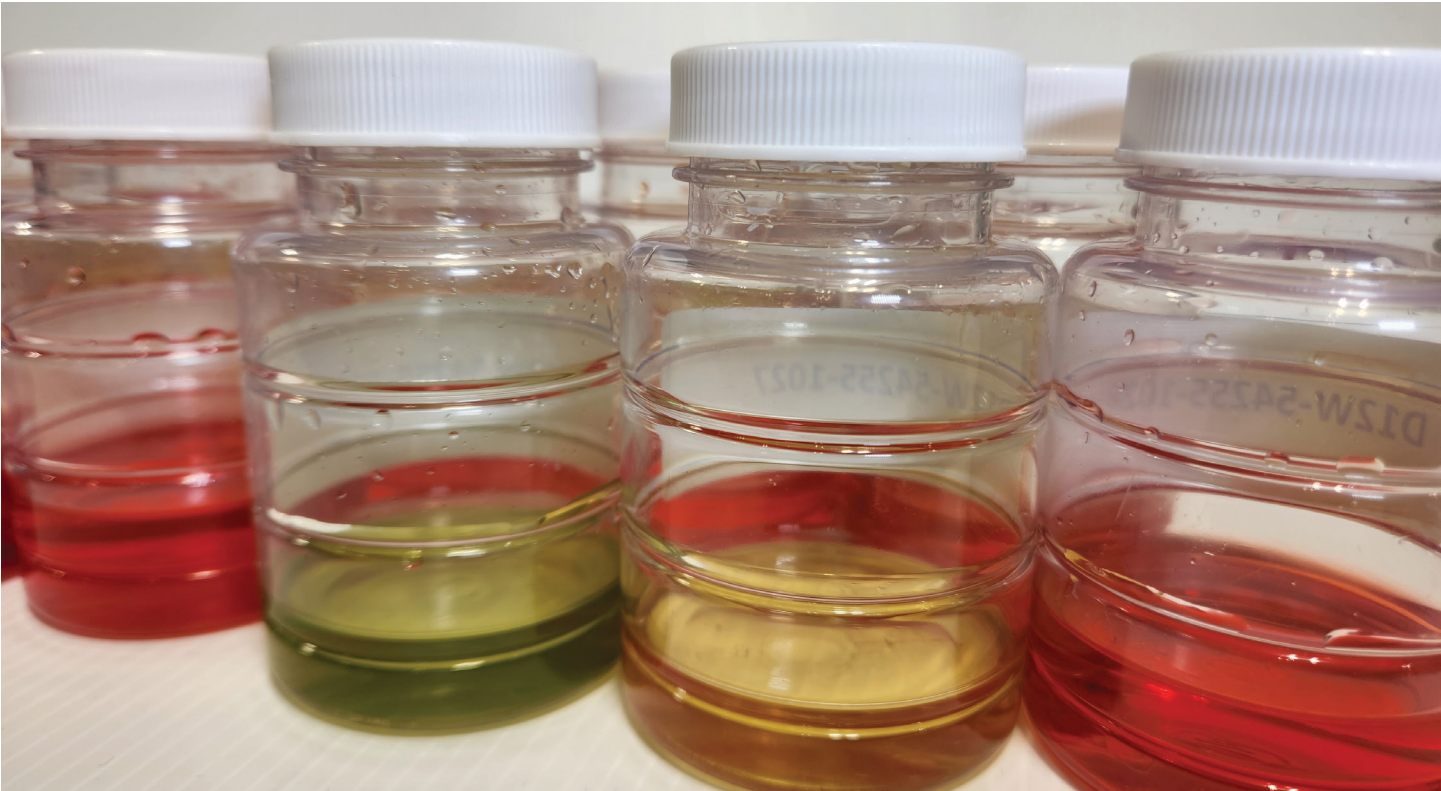


Figure 3. Image of various coolant samples analyzed

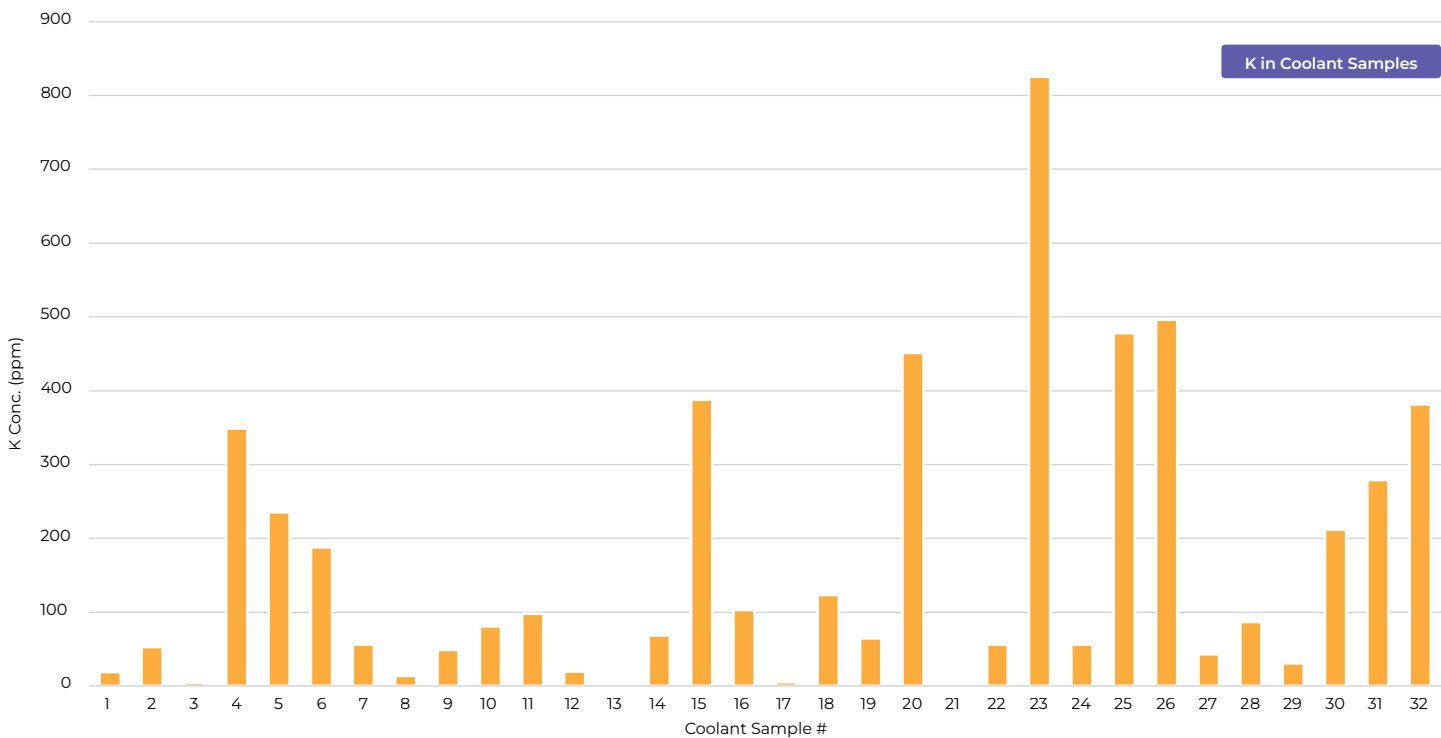


Figure 4. Potassium (K) concentrations measured in coolant samples (ppm)

System Stability

The coolant method was also evaluated for its stability, as many laboratories need to analyze a large number of coolant samples in a single session. Figure 5 displays the QC Sample stability obtained across the over 5-hour sample analysis, with all the recovery values falling within the $\pm 5\%$ control window required in ASTM Method D-6130¹.

Equally as important is the recovery of the Co internal standard during the analysis of these in-service coolants (Figure 6). This graph demonstrates the importance of the internal standard to correct for the different physical and chemical properties of the coolants.

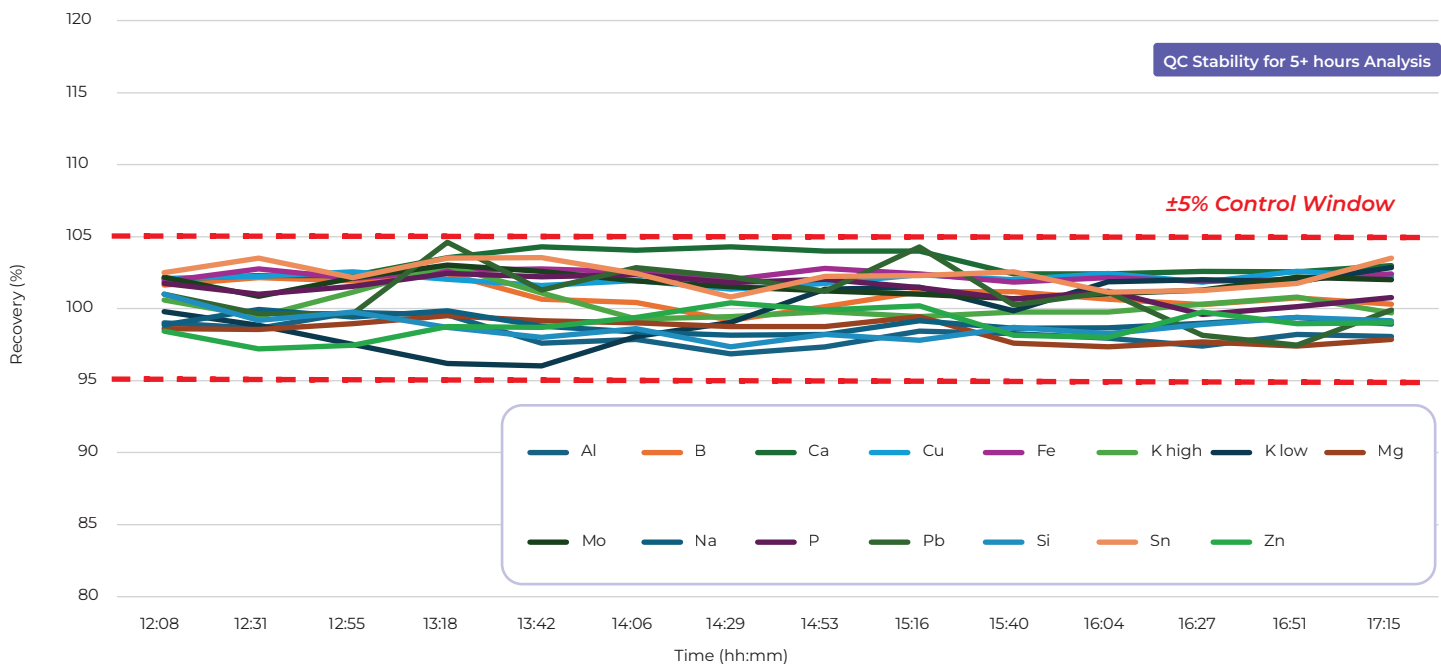


Figure 5. Stability of quality control results analyzed during 5+ hour coolant sample analysis

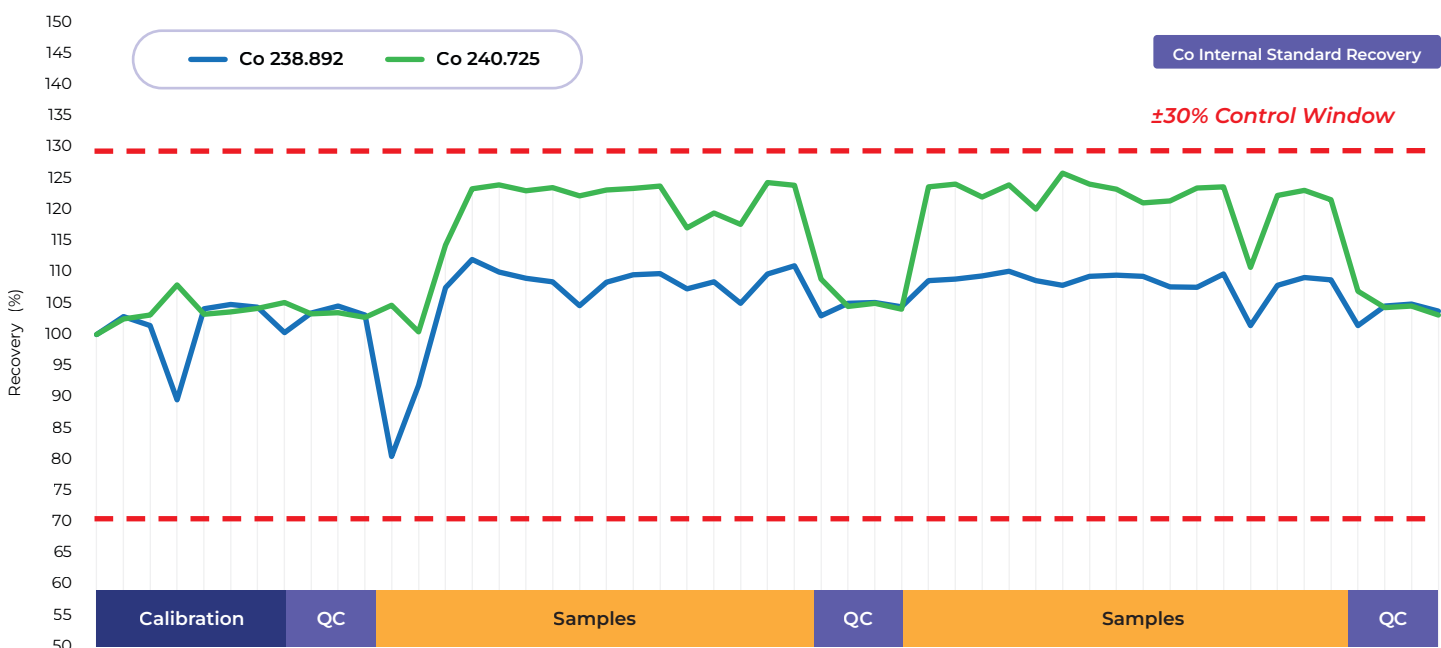


Figure 6. Internal Standard stability observed during coolant sample analysis

Detection Limits

The detection capability of the MICAP in coolants was determined in the glycol matrix utilized for standard preparation. Ten blanks were measured under the method conditions and the detection limits (DL) calculated by:

DL = 3x (Std Dev of 10 blank measurements)

The values obtained are shown in Table 4. These values are presented both as measured and in the actual coolants (reflecting the 10x dilution).

Experimental Conditions

In-service coolant samples often present washout challenges due to their varying composition. The ESI SampleSense FAST automation was primarily selected to help address these challenges. With its automated valve sampling, samples are loaded simultaneously into the valve while the MICAP sample introduction system is rinsed. Additionally, more viscous samples (that load slower) are actively monitored during the loading process. Improved analyte washout is a result of using this technology, resulting in faster sample throughput for quicker analysis times. Figure 7 demonstrates the highly effective sample washout to blank levels immediately after analysis of the high standard levels

The sample throughput rate obtained with a basic autosampler was determined to be 2 min 5 sec per sample. Utilizing the ESI SampleSense FAST automation significantly reduced this to 1 min 11 sec per sample, providing a 43% increase in sample throughput along with enhanced washout performance.

Table 4. Detection limits in glycol matrix

Analyte Wavelength (nm)	DL In Solution (ppm)	DL In Coolant (ppm)
Al 396.152	0.006	0.065
B 249.677	0.061	0.607
Ca 396.847	0.000	0.000
Cu 324.754	0.002	0.021
Fe 259.940	0.008	0.083
K 766.490	0.011	0.110
Mg 280.270	0.002	0.024
Mo 317.034	0.032	0.325
Na 568.819	0.573	5.726
P 213.618	0.573	5.734
Pb 283.305	0.062	0.618
Si 251.611	0.023	0.232
Sn 283.998	0.059	0.595
Zn 213.857	0.022	0.221

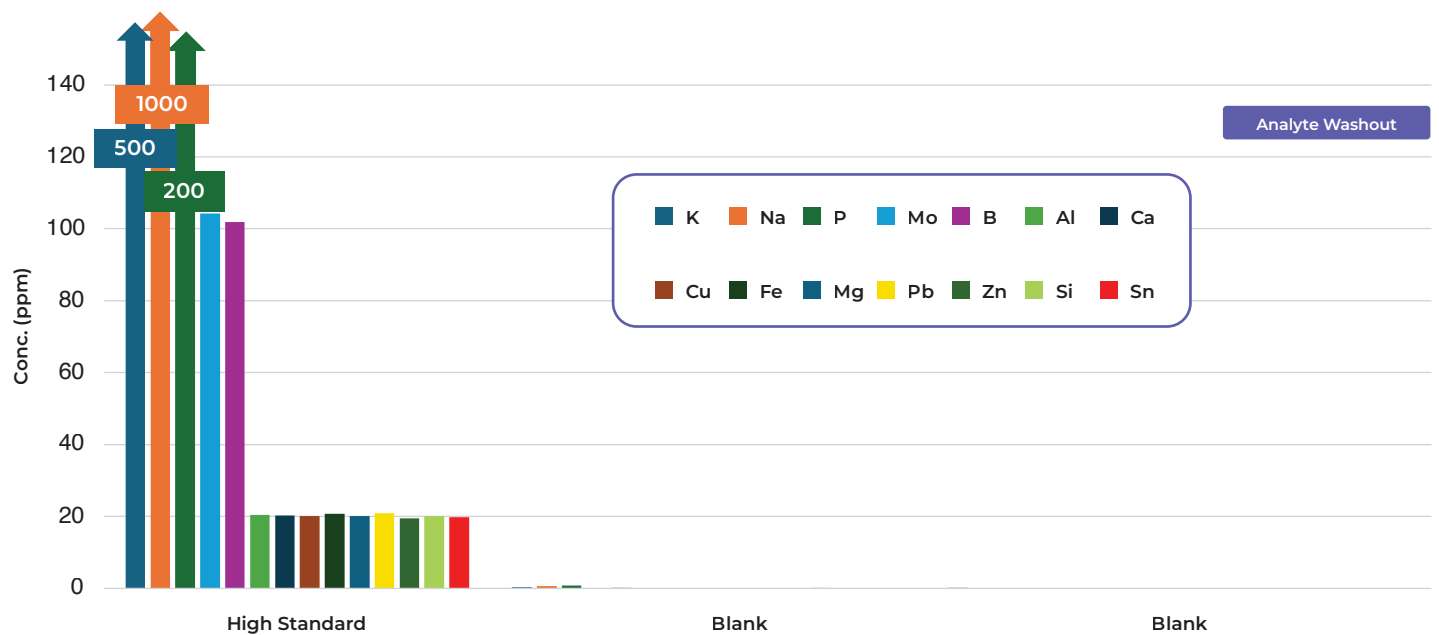


Figure 7. Analyte washout on MICAP with ESI SampleSense FAST. Washout to blank levels immediately after high standard levels. Note 500 ppm K, 1000 ppm Na, & 200 ppm P are off scale in the high standard.

Conclusion

The performance of the MICAP-OES 1000 N2 ICP-OES demonstrates its capability for the high throughput analysis of new and in-service coolants in accordance with ASTM Standard Method D6130-24. The high matrix tolerance of the system coupled with the proper application of internal standard correction delivers stable and accurate monitoring of these critical coolants.

Utilization of the ESI SampleSense *FAST* automated introduction system ensured that the varying viscosity coolant samples were each monitored for the optimal sample loading time, while also enhancing the sample washout process. An average analysis time of 71 seconds/sample was obtained with the ESI automation coupled to the MICAP.

References

1. ASTM D6130-24, Standard Test Method for Determination of Silicon and Other Elements in Engine Coolant by Inductively Coupled Plasma-Atomic Emission Spectroscopy, ASTM International, West Conshohocken, PA, 2024, www.astm.org

Typical Limit of Detection and Resolution

Typical Resolution

1

H

HYDROGEN

2

He

HELIUM

3

Li

19 ppb

BERYLLIUM

4

Be

2 ppb

BERYLLIUM

5

B

8 ppb

BORON

6

C

9 ppb

CARBON

7

N

NITROGEN

8

O

OXYGEN

9

F

FLUORINE

10

Ne

NEON

11

Na

2 ppb

SODIUM

12

Mg

1 ppb

MAGNESIUM

13

Al

6 ppb

ALUMINUM

14

Si

14 ppb

SILICON

15

P

560 ppb

PHOSPHORUS

16

S

SULFUR

17

Cl

CHLORINE

18

Ar

ARGON

19

K

24 ppb

POTASSIUM

20

Ca

1 ppb

CALCIUM

21

Sc

1 ppb

SCANDIUM

22

Ti

7 ppb

TITANIUM

23

V

9 ppb

VANADIUM

24

Cr

8 ppb

CHROMIUM

25

Mn

2 ppb

MANGANESE

26

Fe

12 ppb

IRON

27

Co

21 ppb

COBALT

28

Ni

27 ppb

NICKEL

29

Cu

2 ppb

COPPER

30

Zn

26 ppb

ZINC

31

Ga

402.298nm

GALLIUM

32

Ge

32 ppb

GERMANIUM

33

As

330 ppb

ARSENIC

34

Se

290 ppb

SELENIUM

35

Br

BROMINE

36

Kr

KRYPTON

37

Rb

RUBIDIUM

38

Sr

1 ppb

STRONTIUM

39

Y

1 ppb

YTTORIUM

40

Zr

7 ppb

ZIRCONIUM

41

Nb

24 ppb

NIOBIUM

42

Mo

22 ppb

MOLYBDENUM

43

Tc

TECHNETIUM

44

Ru

12 ppb

RUTHENIUM

45

Rh

322.078nm

RHODIUM

46

Pd

8 ppb

PALLADIUM

47

Ag

10 ppb

SILVER

48

Cd

13 ppb

CADMIUM

49

In

451.131nm

INDIUM

50

Sn

110 ppb

TIN

51

Sb

252.853nm

ANTIMONY

52

Te

238.279nm

TELLURIUM

53

I

14,000 ppb

IODINE

54

Xe

XENON

55

Cs

1 ppb

CAESIUM

56

Ba

1 ppb

BARIUM

57

La

LANTHANUM

58

Ce

CERIUM

59

Pr

PRASEODYMIUM

60

Nd

NEODYMIUM

61

Pm

PM

62

Sm

SAMARIUM

63

Eu

EUROPEUM

64

Gd

GD

65

Tb

TERBBIUM

66

Dy

DYSPROSIUM

67

Ho

HOLMIUM

68

Er

ERBIUM

69

Tm

TM

70

Yb

YBBIUM

71

Lu

LU

72

Hf

67 ppb

HAFNIUM

73

Ta

51 ppb

TANTALUM

74

W

43 ppb

TUNGSTEN

75

Re

40 ppb

RHENIUM

76

Os

84 ppb

OSMIUM

77

Ir

37 ppb

IRIDIUM

78

Pt

34 ppb

PLATINUM

79

Au

16 ppb

GOLD

80

Hg

37 ppb

MERCURY

81

Tl

94 ppb

THALLIUM

82

Pb

65 ppb

LEAD

83

Bi

83 ppb

BISMUTH

84

Po

PO

85

At

ASTATINE

86

Rn

RADON

87

Fr

FRANCIUM

88

Ra

RADIUM

89

Ac

ACTINIUM

90

Th

THORIUM

91

Pa

PA

92

U

URANIUM

93

Np

NEPTUNIUM

94

Pu

PLUTONIUM

95

Am

AMERICIUM

96

Cm

CURIUM

97

Bk

BERKELIUM

98

Cf

CF

99

Es

ESBORIUM

100

Fm

FERMIDIUM

101

Mendelevium

102

No

NOBOLIUM

103

Lr

LR

104

Rf

RUTHERFORDIUM

105

Db

DUBNIUM

106

Sg

SEABORGIUM

107

Bh

BOHRIUM

108

Hs

HASSIUM

109

Mt

METHELIUM

110

Ds

DARMSTADIUM

111

Rg

ROENTGENIUM

112

Cn

COPERNICIUM

113

Nh

NHOLMIUM

114

Fl

FLEEROVIUM

115

Mc

MOSCOWIUM

116

Lv

LIVERMORIUM

117

Ts

TERNESIUM

118

Og

OGANESSON

1

Wavelength (nm)

Resolution (pm)

Typical Resolution

24

428.973nm

5

5

360.534

11

11

11

455.403

14

14

14

588.996

21

21

21

610.364

30

30

30

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