#### **A Novel Approach for Quantitation of Total Iron and Transferrien Bound Iron in** Human serum samples by using Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES)

Mr Dharmesh Patel, Dr. Shanti S Yadav, Dr. Pritesh Contractor, Dr. Ajay Gupta, Ms. Swati Guttikar, Dr. E. Venu Madhav

Veeda Clinical Research Pvt.Ltd.,

Shivalik Plaza, Near I.I.M., Ambawadi, Ahmedabad – 380 015, India

# **Background:**

Intravenous iron therapy is indicated for the treatment of iron deficiency anemia in several patient populations, especially when a large loss of blood occurs, such as in haemodialysis patients with chronic kidney disease, or when oral iron-supplementation is ineffective.

Physiologically<sup>1</sup>, iron–carbohydrate (iron–sucrose, –gluconate or –dextran) therapies replenish depleted iron stores as iron is released from the carbohydrate complex. Once released from the dosage form, iron is taken up by plasma transferrin proteins that circulate and distribute iron throughout the body.

Pharmacokinetic studies of intravenous iron—sucrose are complicated by background cir-culating iron levels as well as the desire to differentiate and independently monitor iron-sucrose and transferrin-bound iron (TBI). Prior to the administration of iron-sucrose, most circulating iron is in the form of TBI; following intravenous iron-sucrose therapy, circulating iron is found both as a component of the dosed formulation and complexed to transferrin.

#### **Method Summary TOI & TBI:** Stability Experiment Details TOI & TBI:

| Analytical Technique   | Inductively coupled Plasma-Optical Emission  |  | ΤΟΙ  | TBI   |
|--|--|--|--|---|
|  | Spectroscopy   | Stability of Extract (SE) at                   | 124 Hours at in Milli-Q -  | 56 Hours at in Milli-Q -                                |
| ICP-OES  | PerkinElmer Optima 8000  | Ambient Temperature                            | water  | water   |
| Auto-sampler<br>Software used  | PerkinElmer S10 Auto-sampler<br>Syngestix software version No 1.0 (for analysis)<br>and WATSON LIMS 7.3 for final regression | Freeze Thaw (FT)                               | 5 Cycles at freezing<br>temperature of -20±5 °C                        | 5 Cycles at freezing<br>temperature of -20±5 °C         |
| Iron (Fe) wavelength<br>Titanium (Ti) wavelength                             | 259.946nm<br>334.940nm   |  | and -78±8 °C   | and -78±8 °C  |
| Nebulizer<br>Spray Chamber   | Gem Cone Low flow nebulizer<br>Cyclonic  | Bench Top (BT)                                 | 07 Hours at ambient<br>temperature                                     | 09 Hours at ambient<br>temperature                      |
| Sample Flow Rate<br>Biological Matrix<br>Internal Standard<br>Quantification | 1.20mL/min<br>Human Serum<br>Titanium Standard for ICP<br>Measured Peak Intensity  | Auto-sampler Re-Injection<br>Reproducibility   | 123 Hours at Ambient<br>Temperature in Milli-Q<br>Water.               | 69 Hours at Ambient<br>Temperature in Milli-Q<br>Water. |
| Regression & Equation<br>Weighting Factor                                    | Linear, y = ax + b<br>1/X2   | Long Term Stability of<br>Drug in Matrix (LTM) | 139 Days at -20±5°C and -<br>78±8°C                                    | 245 Days at -20±5°C and -<br>78±8°C                     |
| Sample Processing Volume<br>Linearity Range                                  | TOITBI0.200mL0.050mL0.500-60.000μg/mL0.500 – 16.000μg/mL   | Batch Size Experiment                          | Total 141 samples<br>including Calibration<br>Curve                    | Total 132 samples including<br>Calibration Curve        |
| Validated LLOQ<br>Validated LLOQ QC<br>Validated LQC                         | 0.500 μg/mL 0.500 μg/mL<br>1.224 μg/mL 1.970 μg/mL<br>2.224 μg/mL 2.970 μg/mL  | Dilution Intogrity (DI)                        | 10 fold, DQC:<br>167.397µg/mL  | 10 fold, DQC: 87.390µg/mL                               |
| Validated MQC<br>Validated HQC<br>Validated AUL QC                           | 20.724 μg/mL 6.470 μg/mL<br>50.724 μg/mL 13.470 μg/mL<br>167.397 μg/mL 87.390 μg/mL  | Dilution Integrity (DI)                        | Amended endogenous concentration, (Dilution medium used - human Serum) |   |
| Validated ULOQ   | 60.000 μg/mL 16.000 μg/mL  |  |  |   |

#### **Purpose:**

To Develop and validate the reliable method for determination of Total Iron (TOI) and Transferrin-Bound Iron (TBI) in Human Serum Samples by using Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES). The purpose of this study was to develop a robust and accurate method for the analysis of Total Iron (TOI) and Transferrin-Bound Iron (TBI) to generate pharmacokinetic data as a component of bioequivalence studies for

| <b>Why ICP OES (Inductively Coupled Plasma O</b> | ptical |
|--|--------|
| Emission Spectroscopy):                          |        |

For decades ICP has been used in support of Environmental Protection Agency analysis and has more recently been applied for use in the pharmaceutical industry.

Increasingly<sup>2</sup>, ICP–MS and ICP–OES methods of analysis are being used to quantify the concentration of elements contained in pharmaceutical compounds and excipients used in nonclinical and clinical studies and in forensic investigations. Validation of methods for quantification of elements in biomatrices such as whole blood, serum, plasma, urine may be performed following GLP predicate rules and bioanalytical guidances, as long as the amount of endogenous element in the matrix is determined during method development and properly accounted for during method validation.



Elemental analysis plays a role in Bio-analysis when the element is derived from the pharmaceutical being administered. Some examples of these applications are presented below:

**Platinum (Pt)** is a component of cisplatin, carboplatin and oxiplatin: all used for oncology treatments

Iron (Fe) from Iron Sucrose, Ferumoxytol, Ferric Carboxymaltose, Sodium Ferric Gluconate and Iron Dextran

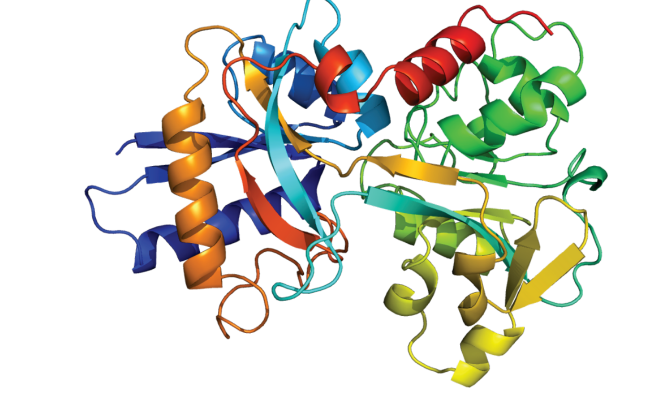
#### **Extraction Cartridges Efficiency Experiment:**

- Formulations (Iron Sucrose 100mg/5mL) spiked QCs were prepared at Higher (HQC) and Lower (LQC) QCs level to demonstrate the efficiency of extraction cartridges. Final spiked concentration of formulation was 60.0µg/mL in blank serum matrix.
  - These formulation spiked HQCs and LQCs (six replicates) each) were processed and evaluated for its acceptance. Results are tabulated below.

|                 | VNF LQC | VNF HQC |
|-----------------|---------|---------|
| Precision (%CV) | 2.09    | 2.55    |
| % Mean Accuracy | 105.75  | 103.68  |

Acceptance Criteria: %CV: within 15.0% and % Mean Accuracy: within ±15.00%.

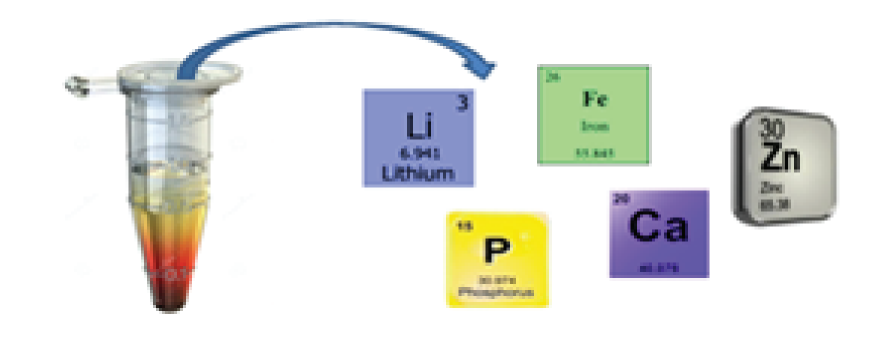
Intra – Inter Precision and Accuracy TOI & TBI:

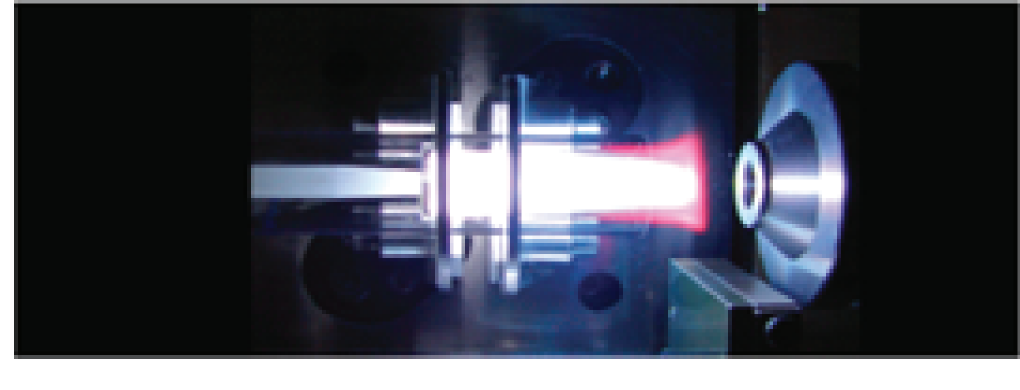


Zinc (Zn) acetate for the treatment of as maintenance treatment in Wilson's disease

Lithium (Li) Carbonate indicated in the treatment of manic episodes of manic-depressive illness.

Potassium (K) chloride For the treatment of patients with hypokalaemia





## **Methodology:**

An exclusive Emission Spectroscopy method was used to reliably measure total iron and TBI following iron-sucrose administration. A validation was conducted in accordance with the method validation protocol in compliance with the regulatory requirements.

Due to its Endogenous presence; analyte free matrix is difficult to obtain. Also, the Endogenous presence varies in individuals. During method development, the measure Intensity of Endogenous drug present in serum matrix (obtain in Serum Blank samples) is subtracted from the calibration curve standards to construct the calibration curve and validate the curve.

Solid-phase sample processing allowed the measurement of TBI. Circulating iron-sucrose could then be calculated as the difference between total iron and TBI. For the analysis of total iron (the sum of free iron, TBI and iron-sucrose) extensive method development was not required.

| ΤΟΙ                              | Precision        |                  |                  | % Bias     |               |          |
|----------------------------------|------------------|------------------|------------------|------------|---------------|----------|
|                                  | RUN 1            | RUN 2            | RUN 3            | RUN 1      | RUN 2         | RUN 3    |
| Intra batch precision and % Bias | < 2.22           | <u>&lt;</u> 3.12 | <u>&lt;</u> 2.05 | -4.65 to - | -1.39 to 1.88 | -2.04 to |
| (Accuracy) (HQC, MQC and LQC)    | $\geq$ Z.ZZ      |                  |                  | 2.79       |               | 0.40     |
| Intra batch precision and % Bias | 1.62             | 1.55             | 2.53             | -4.25      | -4.90         | 3.43     |
| (Accuracy) (LLOQ QC)             | 1.02             | 1.55             | 2.55             | -4.25      | -4.90         | 5.45     |
| Inter batch precision and % Bias | <u>&lt;</u> 3.30 |                  | -2.69 to -0.94   |            |               |          |
| (Accuracy) (HQC, MQC and LQC)    |                  |                  |                  |            |               |          |
| Inter batch precision and % Bias |                  | 1 11             |                  | -1.88      |               |          |
| (Accuracy) (LLOQ QC)             | 4.41             |                  |                  | -1.00      |               |          |

| TBI   | Precision |        |               | % Bias       |               |              |
|---|-----------|--------|---------------|--------------|---------------|--------------|
| IDI   | RUN 1     | RUN 2  | RUN 3         | RUN 1        | RUN 2         | RUN 3        |
| Intra batch precision and % Bias<br>(Accuracy) (HQC, MQC and LQC) | < 3.93    | < 6.07 | < 5.55        | 0.30 to 3.79 | -1.82 to 2.44 | 0.13 to 3.36 |
| Intra batch precision and % Bias<br>(Accuracy) (LLOQ QC)          | 2.27      | 3.28   | 3.72          | 2.99         | -0.91         | -0.30        |
| Inter batch precision and % Bias (Accuracy) (HQC, MQC and LQC)    | < 4.77    |        | -0.47 to 2.73 |              |               |              |
| Inter batch precision and % Bias<br>(Accuracy) (LLOQ QC)          | 3.43      |        | 0.61          |              |               |              |

# **Past Study Experience:**

| STUDY 01                     | ΤΟΙ    | TBI    |
|------------------------------|--------|--------|
| Total No. of Sample Analyzed | 6034   | 6034   |
| Repeat Samples (%)           | 5.30%  | 6.56%  |
| ISR Acceptance (%)           | 81.85% | 86.14% |

#### **Transferrin:**

The predominant iron-binding plasma glycoprotein regulating and distributing circulating physiological levels of iron.

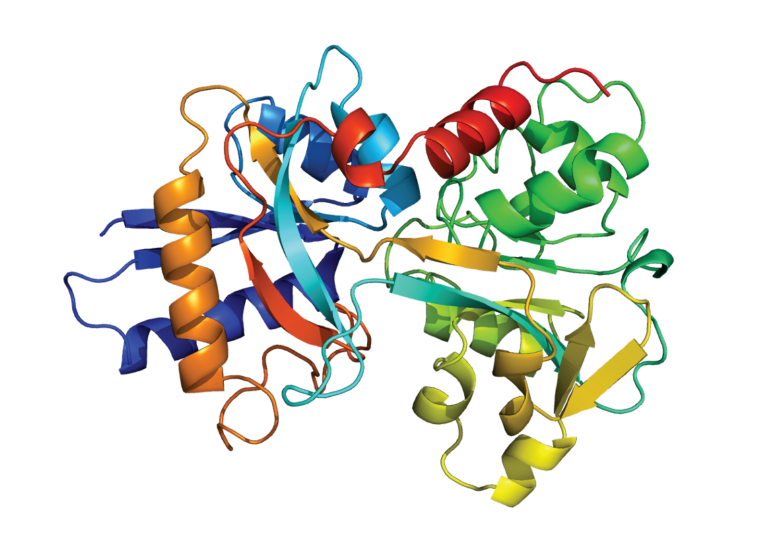
The measurement of TBI requires separating and excluding free iron and drug-derived iron from TBI. Specifically, we encountered difficulty excluding free and sucrose-bound iron for the determination of TBI. SPE columns were evaluated for their ability to remove free and sucrose-bound iron from serum. Method development concludes polyimine extraction cartridges are accurate for determination of TBI.

# **Sample Preparation TOI & TBI:**

Calibration curve standards, Quality control samples and unknown samples (0.200mL for TOI and 0.050mL for TBI) were filled with internal standard (0.100mL) of Titanium and finally diluted with milli-q water as per pre-define procedure.

The measurement of TBI requires separating and excluding free iron and drug-derived iron from TBI. Hence, TBI is separated by applying Solid-phase extraction with the usage of polyimine extraction cartridges. These cartridges were evaluated for their ability to remove free and sucrose bound iron from serum during method development and validation also.

Arrange the final prepared samples in auto-sampler and acquired by applying pre-define equipment parameters.



| STUDY 02                     | ΤΟΙ    | TBI    |  |
|------------------------------|--------|--------|--|
| Total No. of Sample Analyzed | 2211   | 2211   |  |
| Repeat Samples (%)           | 5.29 % | 8.14 % |  |
| ISR Acceptance (%)           | 96.09% | 94.78% |  |

## **Conclusion:**

Two separate Emission Spectroscopy assay has been developed to measure Total Iron (TOI) and Transferrin-Bound Iron (TBI) levels in human serum. Polyimine SPE columns effectively removed free iron and iron-sucrose from serum. Accuracy, precision and incurred sample reanalysis results indicate that the assay is reproducible and robust. The procedures have been applied to clinical analysis of pharmacokinetic studies of generic iv. iron formulations of iron-sucrose.

## **Reference:**

- 1. Melissa M et.al, Analysis of total and transferrin-bound iron in human serum for pharmacokinetic studies of iron–sucrose formulations, Bioanalysis (2011) 3(16), 1837–1846
- 2. Jennifer A et.al, Technical aspects of inductively coupled plasma bioanalysis techniques, Bioanalysis (2013) 5(15), 1831–1841