Interactive comment on “Method development for arsenic analysis by modification in spectrophotometric technique” by M. A. Tahir et al.

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General Comments
This paper describes development of a reliable and cost effective analytical method that can potentially be used in the field to measure total arsenic to less than the regulatory limit of 10 \( \mu \)g/L. The sophisticated and high tech analytical instruments such as, Inductive Coupled Plasma – Mass Spectrometry (ICP MS), Inductive Coupled Plasma – Atomic Emission Spectroscopy (ICP – AES), etc. are capable to achieving very low level arsenic (<1 \( \mu \)g/L). However, the authors concluded that these instruments are expensive and not so easily available. Keeping that objective in mind, the authors made an attempt to develop a low level arsenic measurement process by modification in spectrophotometric technique.

This is an interesting paper. At present, research is being conducted pertaining to measurement of arsenic in the field. If this research concept can be applied to the real world, this would be a real contribution of this work. From the perspective of global arsenic problem, this paper will potentially contribute to the Environmental and Sanitary Engineering Society. The authors have presented the information in a simple and common language such that any engineering practitioners, water suppliers, and operators can understand the method. However, there are some specific points of criticism.

Specific Comments

• The interferences due to phosphate, and chloride on arsenate were not discussed at all. Because of the similar chemistry between arsenate and phosphate, I would assume significant interference due to phosphate. Based on my past experience, I had faced this problem. I’ll appreciate, if the authors clarify this important point

• Page 136 line 23 to 25: Please note that under strongly reduced condition (Negative ORP) arsenic exists as AsH3); under moderately reduced environment \( [E_h = 0.1 \text{ V to } 0.55 \text{ V}] \) arsenic exists as Arsenic (III); under oxidizing environment \( [E_h = 0.60 \text{ V and up}] \) arsenic exists as Arsenic (V) \[ \text{Ref. Arsenic Chemistry: } E_h – \text{pH Diagram; Ferguson and Garvis, 1972} \]

• In Table 3, 10 out of the 15 data point obtained from the spectrometer analyses are slightly higher than those obtained from the AAS. Is there any scientific
reasons?

In conclusion, I wish to complement the authors on a properly presented appropriated topic.