A Method to Study the Effects of Chemical and Biological Reduction of Molybdate to Molybdenum Blue in Bacteria

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Abstract: In this research, we modify a previously developed assay for the quantification molybdenum blue to determine whether inhibitors to molybdate reduction in bacteria inhibits cellular reduction or inhibit the chemical formation of one of the intermediate of molybdenum blue, phosphomolybdate. We manage to prove that inhibition of molybdate reduction by phosphate and arsenate is at the level of phosphomolybdate and not cellular. We also prove that mercury is a physiological inhibitor to molybdate reduction. We suggest the use of this method to assess the effect of inhibitors and activators to molybdate reduction in bacteria.

Key words: Molybdate reduction, E. cloacae strain 48, molybdenum blue, inhibitors

INTRODUCTION

Molybdate reduction to molybdenum blue is a phenomenon that is more than one hundred years old. According to Levine (1925), the phenomenon was first reported in E. coli by Capaldi and Proskauer (1896). Since then, other reports on molybdate reduction by other bacterium has been reported in approximately in a ten-year interval (Jan, 1939; Marchal and Girard, 1948; Woolfolk and Whiteley, 1962; Bautista and Alexander, 1972). After a silence of nearly 13 years, microbial molybdate reduction resurfaced again in a report on its reduction by E. coli K12 (Campbell et al., 1985). Sugio et al. (1987, 1988) reported the reduction of molybdate into molybdenum blue by Thiobacillus ferroxidans and identified the sulfuric iron oxidoreductase (SFO-Rase) as the enzyme responsible for the reduction. Further studies by Yong et al. (1997) however, showed that molybdate reduction in T. ferroxidans is chemically mediated by ferrous iron added into the media; a known fact of molybdate chemistry (Lee, 1977; Sidgwick, 1984). The efforts of Sugio et al. (1988) were followed by Ghani et al. (1993), who reported that another heterotrophic bacterium, Enterobacter cloacae strain 48 (EC 48) was able to reduce molybdate to molybdenum blue. A new mechanism of molybdate reduction in EC 48 was proposed involving phosphomolybdate as an intermediate between molybdate and molybdenum blue (Shukor et al., 2000). It was also shown that unlike Thiobacillus ferroxidans where the reduction of molybdate to molybdenum blue is probably due to ferrous iron, in EC 48 it is predominantly enzymatic (Shukor et al., 2002) using the modified method of Munch and Ottow (1983). Using phosphomolybdate as a substrate, the molybdenum blue was partially purified and characterized (Shukor et al., 2003).

Previously, we have devised a standard curve to quantify molybdenum blue from EC 48 using 12 MP reduced by ascorbate to molybdenum blue (Shukor et al., 2000). This method could be a testing tool to simulate the effect of various interfering substances (inhibitors or activators) on the production of molybdenum blue from E. cloacae strain 48 and for other biological reduction of molybdate to molybdenum blue. This would prevent misleading analysis of the effect of activators and inhibitors to the activity of the molybdenum-reducing microbes in the future.

MATERIALS AND METHODS

Enterobacter cloacae Strain 48 was originally isolated from Chengkau, Malaysia (Ghani et al., 1993) and was grown on agar plate and in low phosphate (2.9 mM phosphate) media (pH 7.0) containing glucose (1%), (NH₄)₂SO₄ (0.3%), MgSO₄·7H₂O (0.05%), NaCl (0.5%), yeast extract (0.05%), Na₂MoO₄·2H₂O (0.242%) and Na₂HPO₄ (0.05%). All chemicals are of analytical grade. Molybdenum blue is produced in this media but not at

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672
high phosphate media (100 mM phosphate). The only difference between the high and low phosphate media is the phosphate concentration.

**Preparation of ascorbate-reduced molybdenum blue:**
Briefly, 12-phosphorous molybdate or phosphomolybdate (BDH) was prepared in distilled water as a 5 mM stock solution and the pH adjusted to pH 5.0 with HCl. Ascorbic acid was prepared as a 25% solution in distilled water and kept at 4°C for a maximum of one week. One hundred microliters from the 12 MP stock solutions was added to 100 µL ascorbic acid. Suitable volumes of chemicals (phosphate, mercury and arsenate) from stock solutions were added and the final volume adjusted to one milliliter with distilled water. After 12 h of incubation, the absorbance was read at the wavelength of 865 nm.

**Preparation of cellular-reduced molybdenum blue:**
EC 48 culture grown in 250 mL high phosphate media (100 mM phosphate) was used in the inhibitor studies for the cellular reduction of phosphomolybdate since it contained high molybdenum-reducing activity although the media did not turn blue. Cells were harvested after 24 h through centrifugation at 10,000 g. The pellet was washed with distilled water and resuspended in 100 mL low phosphate media minus yeast extract and 10 mL aliquots was added into sterile bijou bottles. Incubation period was for 24 h. Phosphate (Na₂HPO₄·2H₂O) was added into the bottles from a 1 M stock solution (pH 7.0) at a final concentration from 10 to 200 mM. Heavy metals such as mercury (HgCl₂, 2H₂O, JT Baker, Phillipsburg, USA) and arsenate (AsH₃O₄·7H₂O, Fluka, USA) were added into reaction vessels in the same manner as phosphate above. After incubation period has elapsed, the wavelength at OD 865 nm was read. Concentration of inhibitor causing 50% reduction in activity (IC₅₀) was calculated using a one-phase exponential decay model for non-linear regression analysis performed using GraphPad Prism version 4.00 for Windows, GraphPad Software, San Diego California USA, www.graphpad.com. The experiments above were carried out in triplicate.

**RESULTS AND DISCUSSION**

When phosphate was added into EC 48 media, molybdenum blue production from EC 48 decreased significantly. A similar profile was obtained with the ascorbate reduced 12 MP (Fig. 1). The inhibitory effect on the molybdenum reduction in EC 48 is probably due to either the 12 MP complex not forming since molybdate is not converted to the polyions form at neutral pH (pH 7) or (and) through the disruption of the phosphomolybdate complex at high concentration of phosphate ions. Both conditions could occur simultaneously as observed by Glenn and Crane (1956) who reported that molybdenum blue is unstable in neutral pH and high concentration of phosphate (100 mM). Campbell *et al.* (1985) also reported that cellular reduction of phosphomolybdate by *E. coli* K-12 requires an acidic pH (pH 5-6) and high concentration of phosphate exceeding 50 mM is inhibitory to molybdate reduction.

Figure 2 shows that arsenate appears to inhibit Mo-reducing enzyme activity. Arsenate is an inhibitor to oxidative and substrate level phosphorylation.
CONCLUSION

Since the molybdenum blue assay employing ascorbate reduced 12 MP could simulate closely the profile of inhibition of molybdenum blue from EC-48, the assay can be a testing tool for interfering factors such as chemical and physical, inhibition or activation at the cellular or enzymatic level to confirm the effect of inhibitors or activators on molybdate reduction. The good correlation between cellular and chemical inhibition also strengthen the evidence that phosphomolybate is involved during molybdate reduction in EC-48.

REFERENCES


