

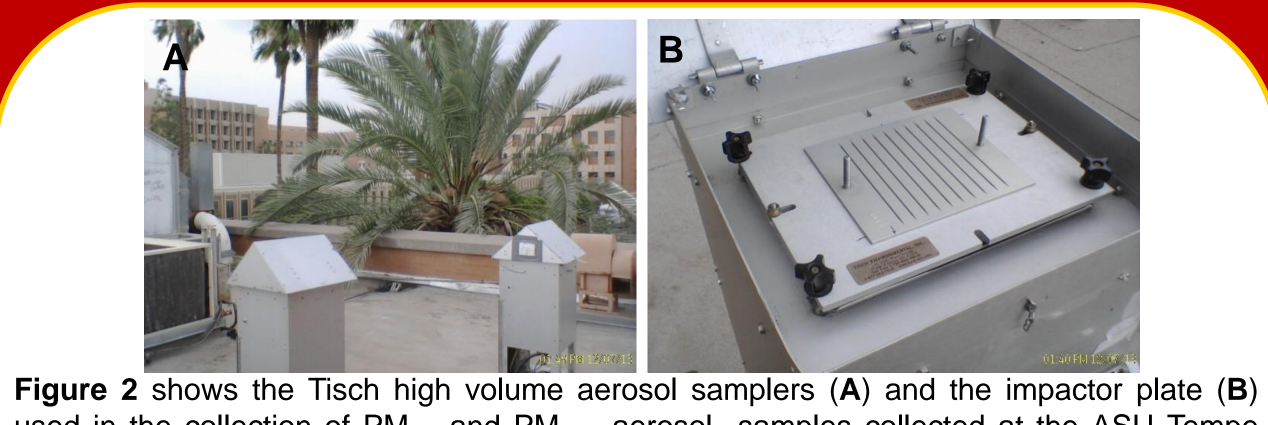
Introduction

During the summer months in Arizona, very intense dust storms, or haboobs, can occur. A haboob formation occurs suddenly when fast moving downdrafts from dissipating thunderstorms hit the ground and, in arid regions, cause the resuspension of dust. This turbulent air mass travels at approximately 30 mph, can be up to 60 miles wide and are an average of 4 000 – 8 000 feet in height. These dust storms can last from minutes to hours and have the potential to alter the aerosol content greatly on short time scales¹. Monsoon-like storms sometimes follow haboobs, which can scavenge the aerosol particles that were brought in by the dust storm. Understanding haboobs is of great importance in the Phoenix area as they can increase particulate matter (PM) and bring an influx of PM material from other locations. Deposition of PM may alter soil and water chemistry in the affected areas. In this work, we chemically characterize haboobs and their effect on the air quality in the Phoenix area. During the summer of 2013, PM_{25} and $PM_{>25}$ aerosol samples were collected on the Arizona State University Tempe Campus. Samples were collected before, during, and after haboobs to determine the time resolved effect of haboobs on PM in the Phoenix area. Samples were analyzed for trace metals by Inductively Coupled Plasma Mass Spectrometry (ICP-MS) and soluble iron content by a ferrozine/UV-Vis method.



Figure 1 shows a July 2011 haboob in Phoenix, AZ.

Experimental



used in the collection of $PM_{2.5}$ and $PM_{>2.5}$ aerosol samples collected at the ASU Tempe Campus on the roof of the Life Sciences A-Wing building.

Tisch high volume (1.13 m³ min⁻¹) aerosol samplers were placed on the roof of the Life Sciences A-Wing building at the ASU Tempe Campus. Cellulose filters were used for samples that would be analyzed for metal concentrations. Single stage impactor plates were used to collect size resolved (PM_{25} and $PM_{>25}$) samples. Samples were collected before, during and after a haboob event. The samplers were run for >1 hour to ensure enough sample mass was obtained.

Trace metals and inorganic species in time resolved haboob samples from **ASU Tempe Campus** Aurelie Marcotte, Jershon Eagar, and Pierre Herckes

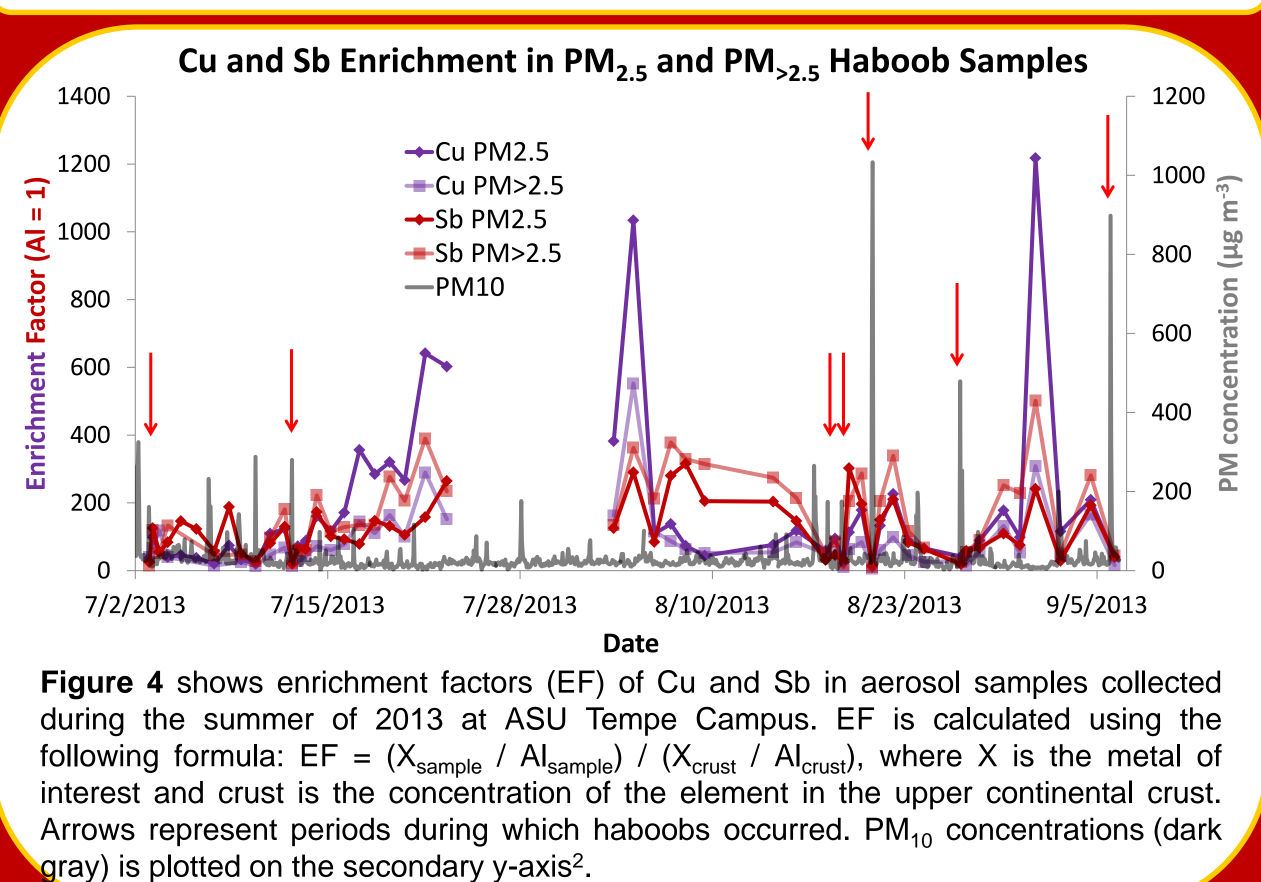
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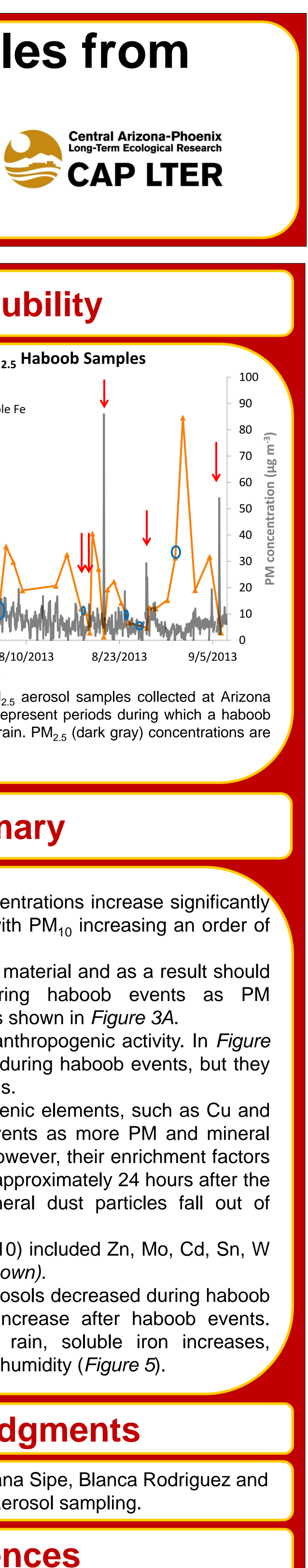
PM Concentrations

Al Concentrations in PM_{>2.5} Haboob Samples 10000 9000 8000 —PM10 7000 6000 5000 4000 2000 1000 7/28/2013 Date Cd Concentrations in PM_{2.5} Haboob Samples ←Cd PM2.5 —PM2.5 7/2/2013 7/15/2013

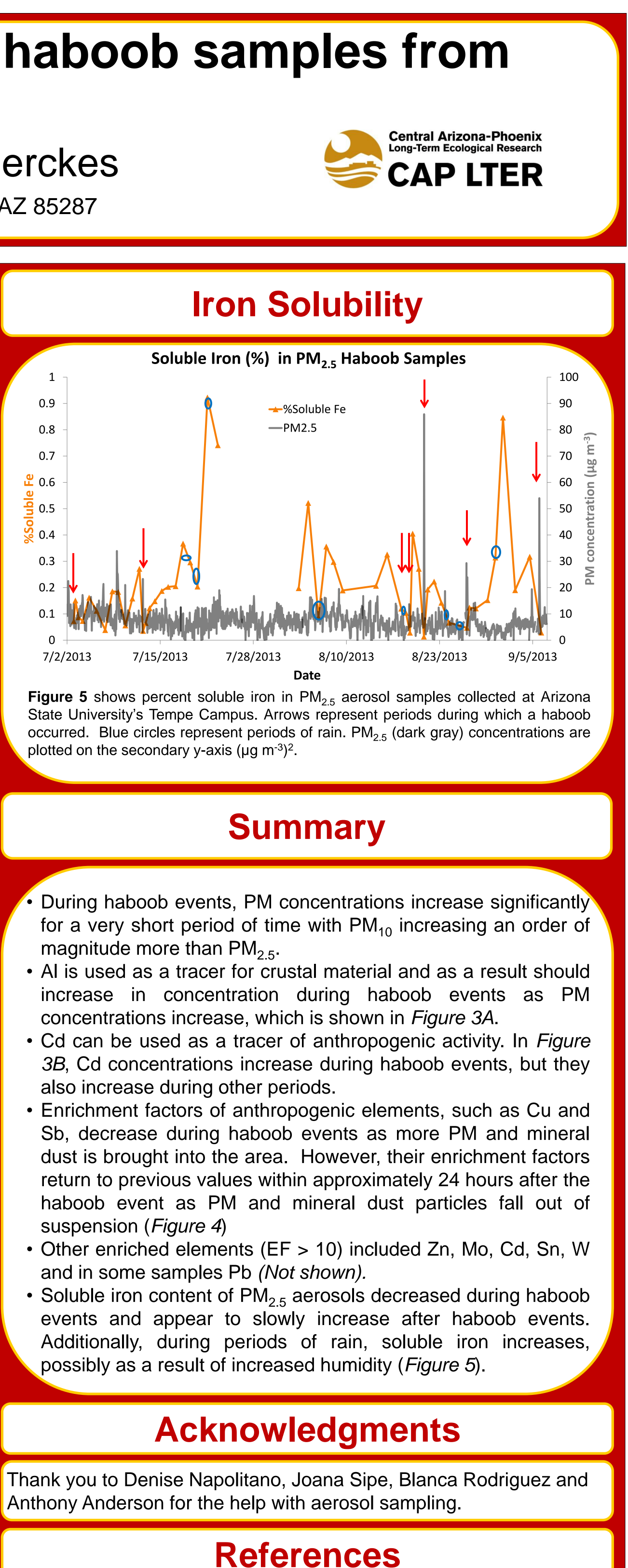
Figure 3 shows particulate matter (PM) concentrations (ng m⁻³) of AI (A; Blue) and Cd (B; Green) in aerosol samples collected during the summer of 2013 at ASU Tempe Campus. Arrows represent periods during which haboobs occurred. PM₁₀ (A) and $PM_{2.5}(B)$ (dark gray) concentrations are plotted on the secondary y-axis (µg m⁻³)².

Enrichment Factors





1000 800 600 400 9/5/2013 100 90 80 70 50 40 30 20 8/23/2013 9/5/2013



¹ Idso, S. B., R. S. Ingram, J. M. Pritchard, 1972: An American Haboob. Bull. Amer. Meteor. Soc., 53, 930–935. ² The Environmental Protection Agency air monitoring site was used to obtain PM_{2.5} and PM₁₀ concentrations.