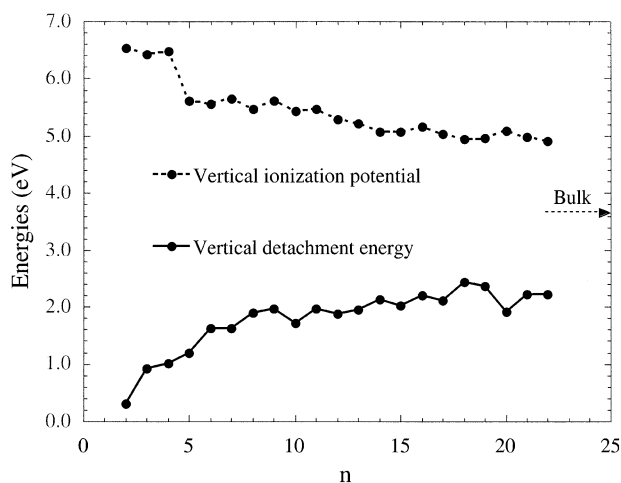


# ADDITIONS AND CORRECTIONS

2002, Volume 106A

**Julius Jellinek\*** and **Paulo H. Acioli** : Magnesium Clusters: Structural and Electronic Properties and the Size-Induced Nonmetal-to-Metal Transition

Page 10924. As a consequence of a technical oversight, the graphs in Figure 9 incurred some unintended changes. The correct figure is given here. The caption of the figure and its discussion in the text are correct and remain unchanged.



**Figure 9.** Vertical ionization potential of the neutral  $Mg_n$  clusters and the vertical electron detachment energy of the anionic  $Mg_n^-$  clusters. The value of the work function of the bulk magnesium is also indicated.

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2003, Volume 107A

**Aly J. Castellanos,\* German Urbina-Villalba, and Máximo García-Sucre** : Mesoscopic Treatment of a Fluid/Liquid Interface. 1. Theory

Pages 875–882. This paper was published in *J. Phys. Chem. A* in the subsection “Molecular Structure, Bonding, Quantum Chemistry, and General Theory” but should have appeared in *J. Phys. Chem. B* in the subsection “Physical Chemistry of Surfaces and Interfaces”.

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**Aly J. Castellanos,\* German Urbina-Villalba, and Máximo García-Sucre** : Mesoscopic Treatment of a Fluid/Liquid Interface. 2. Air/Water Interfacial Tension

Pages 883–887. This paper was published in *J. Phys. Chem. A* in the subsection “Molecular Structure, Bonding, Quantum Chemistry, and General Theory” but should have appeared in *J. Phys. Chem. B* in the subsection “Physical Chemistry of Surfaces and Interfaces”.

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