## **Errata**

In the June 1997 issue of the Journal, in the article entitled, "Characterization of FMR1 Promoter Elements by In Vivo–Footprinting Analysis," by Schwemmle et al.

(60:1354–1362), the ninth author's surname is misspelled; the correct spelling is "Doerfler." The authors regret this error.

In the August 2003 issue of the *Journal*, in the article entitled, "Mutations of von Hippel-Lindau Tumor-Suppressor Gene and Congenital Polycythemia," by Pastore et al. (73:412–419), we reported mutations in the von Hippel-Lindau tumor-suppressor gene as a cause of congenital polycythemia. It was brought to our attention that one of the mutations was mistak-

enly labeled. The 574C→T missense mutation observed in patient 4 results in an amino acid change from proline to serine, not to alanine, as stated; therefore, it should have read: "a previously undescribed 574C→T (P192S) VHL mutation." We thank Andrew Phillip of the Human Genome database for bringing this error to our attention.

In the January 2004 issue of the *Journal*, an error was introduced during production into the article "A Novel Syndrome Combining Thyroid and Neurological Abnormalities Is Associated with Mutations in a Monocarboxylate Transporter Gene," by Dumitrescu et al. (74:168–175). In the original manuscript submission,

the authors included a note acknowledging that Theo Visser's group had made similar findings, which were to be reported at an upcoming meeting. This was mistakenly designated a "Note Added in Proof" in the published version of the paper. The *Journal* regrets this error.