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References and Notes

1. J. W. Lee, C. V. Tevault, T. G. Owens, E. Greenbaum, *Science* **273**, 364 (1996); see also J. M. Olson, *ibid.* **275**, 996 (1997); T. G. Owens, J. W. Lee, C. V. Tevault, E. Greenbaum, *ibid.*, p. 996.
2. E. Greenbaum, J. W. Lee, C. V. Tevault, S. L. Blankinship, L. J. Mets, *Nature* **376**, 438 (1995).

Corrections and Clarifications

In the letter by Charles A. Zeigler in the issue of 27 June (p. 1955), several errors were introduced during editing. The book *Higher Superstition*, referred to in the last line of column one, is by P. R. Gross and N. Levitt only. Reference 1, *The Flight from Science and Reason*, should have identified P. R. Gross, N. Levitt, and M. W. Lewis as "Eds.," and the publication date should have been "1997," not "1994."

In the Table of Contents for the issue of 13 June, the last name of the first author of the report appearing on page 1719, "HSV-TK gene transfer into donor lymphocytes for control of allogeneic graft-versus-leukemia," was spelled incorrectly. It should have been, "Bonini."

An article by Charles Seife about the upgrade of the Arecibo telescope in Puerto Rico (News & Comment, 20 June, p. 1785) erroneously referred to the dish being in the crater of a dead volcano. It is in a limestone sinkhole.

In Jocelyn Kaiser's News & Comment article "When a habitat is not a home" (13 June, p. 1636), the caption for the photographs of a red-cockaded woodpecker and a pine forest on page 1636 should not have read, "[a] rare red-cockaded woodpecker and the loblolly pine habitat it prefers," but rather, "[a] rare red-cockaded woodpecker and the loblolly pine, one of the habitats it prefers."

In reference 27 (p. 1533) of the report "The source of green light emission determined from a heavy-ion storage ring experiment" by D. Kella *et al.* (6 June, p. 1530), the year was incorrect. The reference should have read, "S. L. Guberman, paper presented at the American Physical Society Meeting, Washington, DC, 18 to 21 April 1997."

The ScienceScope item "Lawmakers target biosphere program" (30 May, p. 1323) mischaracterized a bill in the U.S. House of Representatives concerning the U.S. Man in the Bio-

sphere Program (USMAB). The proposed legislation would kill the program only if Congress failed to authorize USMAB and would remove existing biosphere reserves only if each did not obtain congressional approval within 3 years.

In the report "Burst conditions of explosive volcanic eruptions recorded on microbarographs" by M. M. Morrissey and B. A. Chouet (28 Feb., p. 1290), the label on curve A in figure 1 (p. 1291), which read "5 kPa", was incorrect. It should have read, "0.8 kPa".

The end of note 11 (p. 539) in the report "DCP-1, a *Drosophila* cell death protease essential for development" by Z. Song *et al.* (24 Jan., p. 536) should have stated, "The GenBank accession number for *dcp-1* is AF001464."

Letters to the Editor

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