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References

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2. D. R. Cox, L. B. Epstein, C. J. Epstein, *Proc. Natl. Acad. Sci. U.S.A.* **77**, 2168 (1980).

Corrections and Clarifications

In the editorial "In transition" by Barbara Jasny (20 Oct., p. 359), the genus of the organism *Mycoplasma genitalium* was inadvertently given as "Mycobacterium" in the last line of the first paragraph.

In the first report listed in the Table of Contents in the issue of 13 October (p. 211), the name of author C. M. Lieber was misspelled.

The URL address listed for Pete Goldie in his letter of 13 October (p. 218) was incorrect. It should have read, "<http://lbin.com>"

In the Research News article "Designer tissues take hold" by Robert F. Service (13 Oct., p. 230), Katherine Tweden's name was spelled incorrectly. Also, her experimental results were completed in 3 weeks, not 33.

In Paul Selvin's article "The future university: Leaner and meaner?" (Careers '95: The future

of the Ph.D., 6 Oct., p. 135) it was reported that the University of Michigan's Institute for Mental Health Research was closed during the 1980s. In fact, the institute is now celebrating its 40th anniversary. The institute that was closed was the Institute for the Study of Mental Retardation and Related Disabilities, which had formerly been the Institute for the Study of Mental Retardation. *Science* regrets the error.

In the Book Reviews section of 13 October, the two photographs at the bottom of page 319 were inadvertently interchanged.

The News article "Wing scales may help beat the heat" by Wade Roush (29 Sept., p. 1816), should have described *Urania fulgens* as a moth, not a butterfly.

Letters to the Editor

Letters may be submitted by e-mail (at science_letters@aaas.org), fax (202-289-7562), or regular mail (*Science*, 1333 H Street, NW, Washington, DC 20005). Letters will not be routinely acknowledged. Full addresses, signatures, and daytime phone numbers should be included. Letters should be brief (300 words or less) and may be edited for reasons of clarity or space. Beginning in October 1995, our previous policy of consulting with all letter authors before publication will be discontinued.

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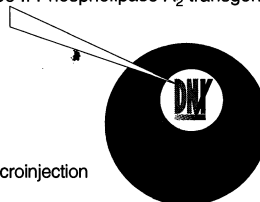
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